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# PATENT APPLICATION

# STREPTOCOCCUS SUIS VACCINES AND DIAGNOSTIC TESTS

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## STREPTOCOCCUS SUIS VACCINES AND DIAGNOSTIC TESTS

[0001] Cross-reference to Related Applications. This application claims priority to, and is a continuation of, International Application No. PCT/NL99/00460, filed on July 19,1999, designating the United States of America, the contents of which are incorporated herein by this reference, the PCT International Patent Application itself claiming priority from European Patent Office Application Serial No. 98202465.5 filed July 22, 1998 and European Patent Office Application Serial No. 98202467.1 filed July 22, 1998.

[0002] Technical Field. The invention relates to *Streptococcus* infections in pigs, vaccines directed against those infections, tests for diagnosing *Streptococcus* infections and bacterial vaccines. More particularly, the invention relates to vaccines directed against *Streptococcus* infections.

#### **Background of the Invention**

[0003] Streptococcus species, of which a large variety cause infections in domestic animals and man, are often grouped according to Lancefield's groups. Typing according to Lancefield occurs on the basis of serological determinants or antigens that are, among others, present in the capsule of the bacterium, and allows for only an approximate determination. Often, bacteria from different groups show cross-reactivity with each other, while other Streptococci [can not]cannot be assigned a group-determinant at all. Within groups, further differentiation is often possible on the basis of serotyping. These serotypes further contribute to the large antigenic variability of Streptococci, a fact that creates an array of difficulties within diagnosis of and vaccination against Streptococcal infections.

[0004] Lancefield group A Streptococcus species (Group A streptococci "GAS", Streptococcus pyogenes) are common in children, causing nasopharyngeal infections and complications thereof. Among animals, cattle are especially susceptible to GAS, and the resulting mastitis.

[0005] Group A streptococci are the etiologic agents of streptococcal pharyngitis and impetigo, two of the most common bacterial infections in children, as well as a variety of less common, but potentially life-threatening, infections including soft tissue infections, bacteremia, and pneumonia. In addition, GAS are uniquely associated with the post-infectious autoimmune syndromes of acute rheumatic fever and post streptococcal glomerulonephritis.

[0006] Several recent reports suggest that the incidence of both serious infections due to GAS and acute rheumatic fever has increased during the past decade, focusing renewed interest on defining the attributes or virulence factors of the organism that may play a role in the pathogenesis of these diseases.

[0007] GAS produce several surface components and extracellular products that may be important in virulence. The major surface protein, M protein, has been studied in the most detail and has been convincingly shown to play a role in both virulence and immunity. Isolates rich in M protein are able to grow in human blood, a property thought to reflect the capacity of M protein to interfere with phagocytosis, and these isolates tend to be virulent in experimental animals.

[0008] Lancefield group B Streptococcus ("GBS") are most often seen in cattle, causing mastitis[,]; however, human infants are susceptible as well, often with fatal consequences. Group B streptococci (GBS) constitute a major cause of bacterial sepsis and meningitis among human neonates born in the United States and Western Europe and are emerging as significant neonatal pathogens in developing countries as well.

[0009] It is estimated that GBS strains are responsible for 10,000 to 15,000 cases of invasive infection in neonates in the United States alone. Despite advances in early diagnosis and treatment, neonatal sepsis due to GBS continues to carry a mortality rate of 15 to 20%. In addition, survivors of GBS meningitis have 30 to 50% incidence of long-term neurologic sequelae. Over the past two decades, increasing recognition of GBS as an important pathogen for human infants has generated renewed interest in defining the bacterial and host factors important in virulence of GBS and in the immune response to GBS infection.

[0010] Particular attention has focused on the capsular polysaccharide as the predominant surface antigen of the organisms. In a modification of the system originally developed by Rebecca Lancefield, GBS strains are serotyped on the basis of antigenic differences

in their capsular polysaccharides and the presence or absence of serologically defined C proteins. While GBS isolated from non[-]human sources often lack a serologically detectable capsule, a large majority of strains associated with neonatal infection belong to one of four major capsular serotypes, la, lb, II or III. The capsular polysaccharide forms the outermost layer around the exterior of the bacterial cell, superficial to the cell wall. The capsule is distinct from the cell wall-associated group B carbohydrate. It has been suggested that the presence of sialic acid, in the capsule of bacteria that causes meningitis, is important for allowing these bacteria to breach the blood-brain barrier. Indeed, in S. agalactiae, sialic acid has been shown to be critical for the virulence function of the type III capsule. The capsule of S. suis serotype is composed of glucose, galactose, N-acetylglucosamine, rhamnose and sialic acid.

[0011] The group B polysaccharide, in contrast to the type-specific capsule, is present on all GBS strains and is the basis for serogrouping the organisms into Lancefield's group B. Early studies by Lancefield and co-workers showed that antibodies raised in rabbits against whole GBS organisms protected mice against challenge with strains of homologous capsular type, demonstrating the central role of the capsular polysaccharide as a protective antigen. Studies in the 1970s by Baker and Kasper demonstrated that cord blood of human infants with type III GBS sepsis uniformly had low or undetectable levels of antibodies directed against the type III capsule, suggesting that a deficiency of anticapsular antibody was a key factor in susceptibility of human neonates to GBS disease.

[0012] Lancefield group C infections, such as those with S. equi, S. zooepidemicus, S. dysgalactiae, and others, are mainly seen in horses, cattle and pigs, but can also cross the species barrier to humans. Lancefield group D (S. bovis) infections are found in all mammals and some birds, sometimes resulting in endocarditis or septicemia.

[0013] Lancefield groups E, G, L, P, U and V (S. porcinus, S[,]. canis, S. dysgalactiae) are found in various hosts, causing neonatal infections, nasopharyngeal infections or mastitis.

[0014] Within Lancefield groups R, S, and T[,] (and with ungrouped types), Streptococcus suis is an important cause of meningitis, septicemia, arthritis and sudden death in young pigs (4, 46). Incidentally, it can also cause meningitis in man (1). S. suis strains are usually identified and classified by their morphological, biochemical and serological characteristics (58, 59,

46). Serological classification is based on the presence of specific antigenic polysaccharides. So far, 35 different serotypes have been described (9, 56, 14). In several European countries, *S. suis* serotype 2 is the most prevalent type isolated from diseased pigs, followed by serotypes 9 and 1. Serological typing of *S. suis* is performed using different types of agglutination tests. In these tests, isolated and biochemically characterized *S. suis* cells are agglutinated with a panel of 35 specific sera. These methods are very laborious and time-consuming.

[0015] Little is known about the pathogenesis of the disease caused by S. suis, let alone about its various serotypes such as type 2. Various bacterial components, such as extracellular and cell-membrane associated proteins, fimbriae, [hemaglutinins] hemagglutinins, and [hemolysis] hemolysin have been suggested as virulence factors (9, 10, 11, 15, 16, 47, 49). However, the precise role of these protein components in the pathogenesis of the disease remains unclear (37). It is well known that the polysaccharide capsule of various Streptococci and other [gram-positive] Gram-positive bacteria plays an important role in pathogenesis (3, 6, 35, 51, 52). The capsule enables these [micro-organisms] microorganisms to resist phagocytosis and is therefore regarded as an important virulence factor. Recently, a role of the capsule of S. suis in the pathogenesis was suggested as well (5). However, the structure, organization and function of the genes responsible for capsule polysaccharide synthesis ([cps]"cps") in S. suis is unknown. Within S. suis, serotype[s] 1 and 2, strains can differ in virulence for pigs (41, 45, 49). Some type 1 and 2 strains are virulent, other strains are not. Because both virulent and non[-]virulent strains of serotype 1 and 2 strains are fully encapsulated, it may even be that the capsule is not a relevant factor required for virulence.

[0016] Attempts to control *S. suis* infections or disease are still hampered by the lack of knowledge about the epidemiology of the disease and the lack of effective vaccines and sensitive diagnostics. It is well known and generally accepted that the polysaccharide capsule of various Streptococci and other gram-positive bacteria plays an important role in pathogenesis. The capsule enables these [micro-organisms] microorganisms to resist phagocytosis and is therefore regarded as an important virulence factor.

[0017] Compared to encapsulated S. suis strains, non-encapsulated S. suis strains are phagocytosed by murine polymorphonuclear leucocytes to a greater degree. Moreover, an

increase in thickness of capsule was noted for *in vivo* grown virulent strains while no increase was observed for avirulent strains. Therefore, these data again demonstrate the role of the capsule in the pathogenesis for *S. suis* as well.

[0018] Ungrouped Streptoccus species, such as S. mutans, causing car[r]ies in humans, S[,]\_ causing mastitis in cattle, and S. pneumonia, causing major infections in humans, and Enterococcus faecilalis and E. faecium, further contribute to the large group of Streptococci.

[0019]Streptococcus pneumoniae (the pneumococcus) is a human pathogen causing invasive diseases, such as pneumonia, bacteremia, and meningitis. Despite the availability of antibiotics, pneumococcal infections remain common and can still be fatal, especially in high-risk groups, such as young children and elderly people. Particularly in developing countries, many children under the age of five years die each year from pneumococcal pneumonia. S. pneumoniae is also the leading cause of otitis media and sinusitis. These infections are less serious, but nevertheless incur substantial medical costs, especially when leading to complications, such as permanent deafness. The normal ecological niche of the pneumococcus is the nasopharynx of man. The entire human population is colonized by the pneumococcus at one time or another, and at a given time, up to 60% of individuals may be carriers. Nasopharyngeal carriage of pneumococci by man is often accompanied by the development of protection against infection by the same serotype. Most infections do not occur after prolonged carriage but follow exposure to recently acquired strains. Many bacteria contain surface polysaccharides that act as a protective layer against the environment. Surface polysaccharides of pathogenic bacteria usually make the bacteria resistant to the defense mechanisms of the host, for example, the lytic action of serum or In this respect, the serotype-specific capsular polysaccharide ("CP") of phagocytosis. Streptococcus pneumoniae, is an important virulence factor. Unencapsulated strains are avirulent. and antibodies directed against the CP are protective. Protection is serotype specific; each serotype has its own, specific CP structure. Ninety different capsular serotypes have been identified. Currently, CPs of 23 serotypes are included in a vaccine.

[0020] Vaccines directed against *Streptococcus* infections typically aim to utilize an immune response directed against the polysaccharide capsule of the various *Streptococcus species*, especially since the capsule is considered a primary virulence factor for these bacteria. During

infection, the capsule provides resistance against phagocytosis and thus protects the bacteria from the immune system of the host, and from elimination by macrophages and neutrophils.

[0021] The capsule particularly confers the bacterium resistance to complement-mediated opsonophagocytosis. In addition, some bacteria express capsular polysaccharides (CPs) that mimic host molecules, thereby avoiding the immune system of the host. Also, even when the bacteria have been phagocytosed, intracellular killing is hampered by the presence of a capsule.

[0022] It is generally thought that the bacterium will [get] be recognized by the immune system through the anticapsular-antibodies or serum-factors bound to its capsule, and will, through opsonization, [get] be phagocytosed and killed only when the host has antibodies or other serum factors directed against capsule antigens.

[0023] However, these antibodies are serotype-specific, and will often only confer protection against only one of the many serotypes known within a group of *Streptococci*.

[0024] For example, current commercially available *S. suis* vaccines, which are generally based on whole-cell-bacterial preparations, or on capsule-enriched fractions of *S. suis*, confer only limited protection against heterologous strains. Also, the current pneumococcal vaccine, [that]which was licensed in the United states in 1983, consists of purified CPs of 23 pneumococcal serotypes whereas at least 90 CP types exist.

[0025] The composition of this pneumococcal vaccine was based on the frequency of the occurrence of disease isolates in the US and cross-reactivity between various serotypes. Although this vaccine protects healthy adults against infections caused by serotypes included in the vaccine, it fails to raise a protective immune response in infants younger than 18 months and it is less effective in elderly people. In addition, the vaccine confers only limited protection in patients with immunodeficiencies and hematology malignancies.

[0026] Thus, improved vaccines are needed against *Streptococcus* infections. Much attention is directed toward producing CP vaccines by producing the relevant polysaccharides via chemical or recombinant means. However, chemical synthesis of polysaccharides is costly, and capsular polysaccharide synthesis by recombinant means necessitates knowledge about the relevant genes, which is not always available, and needs to be determined for every relevant serotype.

#### Disclosure of the Invention

[0027] The invention provides an isolated or recombinant nucleic acid encoding a capsular (cps) gene cluster of Streptococcus suis. Biosynthesis of capsule polysaccharides has generally been studied in a number of Gram-positive and Gram-negative bacteria (32). In Gramnegative bacteria, but also in a number of [g]Gram-positive bacteria, genes which are involved in the biosynthesis of polysaccharides are clustered at a single locus.

[0028] Streptococcus suis capsular genes, as provided by the invention, show a common genetic organization involving three distinct regions. The central region is serotype specific and encodes enzymes responsible for the synthesis and polymerization of the polysaccharides. The central region is flanked by two regions conserved in Streptococcus suis which encode proteins for common functions, such as transport of the polysaccharide across the cellular membrane. However, between species, only low homologies exist, hampering easy comparison and detection of seemingly similar genes. Knowing the nucleic acid encoding the flanking regions allows type-specific determination of nucleic acid of the central region of Streptococcus suis serotypes, as, for example, described herein.

[0029] The invention provides an isolated or recombinant nucleic acid encoding a capsular gene cluster of *Streptococcus suis* or a gene or gene fragment derived thereof. Such a nucleic acid is, for example, provided by hybridizing chromosomal DNA derived from any one of the *Streptococcus suis* serotypes to a nucleic acid encoding a gene derived from a *Streptococcus suis* serotype 1, 2 or 9 capsular gene cluster, as provided by the invention (*see* for example, Tables 4 and 5) and cloning of (type-specific) genes as, for example, described herein. At least 14 open reading frames are identified. Most of the genes belong to a single transcriptional unit, identifying a co[-]ordinate control of these genes[, they,]. The genes and the enzymes and proteins they encode, act in concert to provide the capsule with the relevant polysaccharides.

[0030] The invention provides *cps* genes and proteins encoded thereof involved in regulation (CpsA), chain length determination (CpsB, C), export (CpsC) and biosynthesis (CpsE, F, G, H, J, K). Although, at first glance, the overall organization seemed to be similar to that of the *cps* and *eps* gene[,] clusters of a number of Gram-positive bacteria (19, 32, 42), overall

homologies are low (see, table 3). The region involved in biosynthesis is located at the center of the gene cluster and is flanked by two regions containing genes with more common functions.

[0031] The invention provides an isolated or recombinant nucleic acid encoding a capsular gene cluster of *Streptococcus suis* serotype 2, or a gene or gene fragment derived thereof, preferably as identified in FIG. 3. Genes in this gene cluster are involved in polysaccharide biosynthesis of capsular components and antigens. For a further description of such genes see, for example, Table 2. For example, a cpsA gene is provided functionally encoding regulation of capsular polysaccharide synthesis, whereas cpsB and cpsC are functionally involved in chain[]-in-chain length determination. Other genes, such as cpsD, E, F, G, H, I, J, K and related genes, are involved in polysaccharide synthesis, functioning, for example, as glucosyl[-] or glycosyltransferase. The cpsF, G, H, I, J genes encode more type-specific proteins than the flanking genes which are found more-or-less conserved throughout the species and can serve as a base for selection of primers or probes in PCR-amplification or cross-[hybridisation]hybridization experiments for subsequent cloning.

[0032] The invention further provides an isolated or recombinant nucleic acid encoding a capsular gene cluster of *Streptococcus suis* serotype 1 or a gene or gene fragment derived thereof, preferably as identified in FIG. 4.

[0033] In addition, the invention provides an isolated or recombinant nucleic acid encoding a capsular gene cluster of *Streptococcus suis* serotype 9 or a gene or gene fragment derived thereof, preferably as identified in FIG. 5.

[0034] Furthermore, the invention provides, for example, a fragment of the *cps* locus or parts thereof, involved in the capsular polysaccharide biosynthesis, of *S. suis*, exemplified herein for serotypes 1, 2 or 9, and allows easy identification or detection of related fragments derived of other serotypes of *S. suis*.

[0035] The invention provides a nucleic acid probe or primer derived from a nucleic acid according to the invention allowing species or serotype specific detection of *Streptococcus suis*. Such a probe or primer (used interchangeably herein) is, for example, a DNA, RNA or PNA (peptide nucleic acid) probe hybridizing with capsular nucleic acid as provided by the invention. Species[]-specific detection is provided preferably by selecting a probe or primer sequence from a

species-specific region (e.g. flanking region) whereas serotype[]\_specific detection is provided preferably by selecting a probe or primer sequence from a type-specific region (e.g. central region) of a capsular gene cluster as provided by the invention. Such a probe or primer can be used in a further unmodified form, for example, in cross-hybridization or polymerase-chain reaction (PCR) experiments as, for example, described in the experimental part herein. The invention provides the isolation and molecular characterization of additional type[]\_specific cps genes of S. suis types 1 and 9. In addition, we describe the genetic diversity of the cps loci of serotypes 1, 2 and 9 among the 35 S. suis serotypes known. Type-specific probes are identified. Also, a type-specific PCR, for example, for serotype 9, is provided, being a rapid, reliable and sensitive assay[,] used directly on nasal or tonsillar swabs or other samples of infected or carrier animals.

[0036] The invention also provides a probe or primer according to the invention with at least one reporter molecule. Examples of reporter molecules are manifold and known in the art[,], for example, a reporter molecule can include additional nucleic acid provided with a specific sequence (e.g. oligo-dT) hybridizing to a corresponding sequence [to]in which hybridization can easily be detected, for example, because it has been immobilized to a solid support.

[0037] Yet other reporter molecules include chromophores, e.g. fluorochromes for visual detection, for example, by light microscopy or fluorescent [in situ hybridisation] in situ hybridization ("FISH") techniques, or include an enzyme such as horseradish peroxidase for enzymatic detection, [e.g.] for example, in enzyme-linked assays ("EIA"). Yet other reporter molecules include radioactive compounds for detection in [radiation-based-assays]radiation-based assays.

[0038] In a preferred embodiment of the invention, at least one probe or primer according to the invention is provided (labeled) with a reporter molecule and a quencher molecule, together with <u>an</u> unlabeled probe or primer in a PCR-based test allowing rapid detection of specific hybridization.

[0039] The invention further provides a diagnostic test or test kit including a probe or primer as provided by the invention. Such a test or test kit is, for example, a cross-hybridization test or PCR-based test[,] advantageously used in rapid detection and/or serotyping of Streptococcus suis.

[0040] The invention further provides a protein or fragment thereof encoded by a nucleic acid according to the invention. Examples of such a protein or fragment are[, for example,] proteins described in Table 2. For example, a cpsA protein is provided that functionally encodes regulation of capsular polysaccharide synthesis, whereas cpsB and cpsC are functionally involved in chain[]-in[]-chain length determination. Other proteins or functional fragments thereof, as provided by the invention, such as cpsD, E, F, G, H, I, J, K and related proteins, are involved in polysaccharide biosynthesis, functioning, for example, as glucosyl[-] or glycosyltransferase in polysaccharide biosynthesis of *Streptococcus suis* capsular antigen.

[0041] The invention also provides a method of producing a *Streptococcus suis* capsular antigen including using a protein or functional fragment thereof as provided by the invention, and provides therewith a *Streptococcus suis* capsular antigen obtainable by such a method.

[0042] A comparison of the predicted amino acid sequences of the *cps2* genes with sequences found in the databases allowed the assignment of functions to the open reading frames. The central region contains the type[]-specific glycosyltransferases and the putative polysaccharide polymerase. This region is flanked by two regions encoding for proteins with common functions, such as regulation and transport of polysaccharide across the membrane. Biosynthesis of Streptococcus capsular polysaccharide antigen using a protein or functional fragment thereof is advantageously used in chemo-enzymatic synthesis and the development of vaccines which offer protection against serotype-specific Streptococcal disease, and is also advantageously used in the synthesis and development of multivalent vaccines against Streptococcal infections. Such vaccines elicit ariticapsular antibodies which confer protection.

[0043] Furthermore, the invention provides an acapsular *Streptococcus* mutant for use in a vaccine, a vaccine strain derived thereof and a vaccine derived thereof. Surprisingly, and against the grain of common doctrine, the invention provides use of a *Streptococcus* mutant deficient in capsular expression in a vaccine.

[0044] Acapsular Streptococcus mutants have long been known in the art and can be found in nature. Griffith (J. Hyg. 27:113-159, 1928) demonstrated that pneumococci could be transformed from one type to another. If he injected live rough (acapsular or unencapsulated) type 2 pneumococci into mice, the mice would survive. If, however, he injected the same dose of

live rough type 2 mixed with heat-killed smooth (encapsulated) type 1 into a mouse, the mouse would die, and, from the blood, he could isolate live smooth type 1 pneumococci. At that time, the significance of this transforming principle was not understood. However, understanding came when it was shown that DNA constituted the genetic material responsible for phenotypic changes during transformation.

[0045] Streptococcus mutants deficient in capsular expression are found in several forms. Some are fully deficient and have no capsule at all, others form a deficient capsule, characterized by a mutation in a capsular gene cluster. Deficiency can, for instance, include capsular formation wherein the organization of the capsular material has been rearranged, as, for example, demonstrable by electron microscopy. Yet others have a nearly fully developed capsule which is only deficient in a particular sugar component.

[0046] Now, after much advance of biotechnology and despite the fact that little is still known about the exact localization and sequence of genes involved in capsular synthesis in Streptococci, it is possible to create mutants of Streptococci, for example, by homologous recombination or transposon mutagenesis, which has, for example, been done for GAS (Wessels []et al., PNAS 88:8317-8321, 1991), for GBS (Wessels []et al., PNAS 86: 8983-8987, 1989), for S. suis (Smith, ID-DLO Annual report 1996, page 18-19; Charland []et al., Microbiol. 144:325-332, 1998) and S. pneumoniae (Kolkman []et al., J. Bact. 178:3736-3741, 1996). Such recombinant derived mutants, or isogenic mutants, can easily be compared with the wild-type strains from which they have been derived.

[0047] In a preferred embodiment, the invention provides use of a recombinant-derived Streptococcus mutant deficient in capsular expression in a vaccine. Recombinant techniques useful in producing such mutants are, for example, homologous recombination, transposon [mutagenises]mutagenesis, and others, wherein deletions, insertions or (point)[-mutations] mutations are introduced in the genome. Advantages of using recombinant techniques include the stability of the obtained mutants (especially with homologous recombination and double cross[-]over techniques), and the knowledge about the exact site of the deletion, mutation or insertion.

[0048] In another embodiment, the invention provides a stable mutant deficient in capsular expression obtained, for example, through homologous recombination or [cross over]crossover integration events. Examples of such a mutant can be found herein, for example, mutants lOcpsB or 10cpsEF are stable mutants as provided by the invention.

[0049] The invention also provides a *Streptococcus* vaccine strain and vaccine that has been derived from a *Streptococcus* mutant deficient in capsular expression. In general, the strain or vaccine is applicable within the whole range of Streptococcal infections, including animals or man or with zoonotic infections. It is, of course, now possible to first select a common vaccine strain and derive a *Streptococcus* mutant deficient in capsular expression thereof for the selection of a vaccine strain and use in a vaccine according to the invention.

[0050] In a preferred embodiment, the invention provides use of a *Streptococcus* mutant deficient in capsular expression in a vaccine wherein the *Streptococcus* mutant is selected from the group composed of *Streptococcus* group A, *Streptococcus* group B, *Streptococcus suis* and *Streptococcus pneumoniae*. Herewith the invention provides vaccine strains and vaccines for use with these notoriously heterologous Streptococci, of which a multitude of serotypes exist. With a vaccine, as provided by the invention, that is derived from a specific *Streptococcus* mutant that is deficient in capsular expression, the difficulties relating to lack of heterologous protection can be circumvented since these mutants do not rely on capsular antigens, per se, to induce protection.

[0051] In a preferred embodiment, the vaccine strain is selected for its ability to survive, or even replicate, in an immune-competent host or host cells and thus can persist for a certain period, varying from 1-2 days to more than one or two weeks, in a host, despite its deficient character.

[0052] Although an immunodeficient host will support replication of a wide range of bacteria that are deficient in one or more virulence factors, in general, it is considered a characteristic of pathogenicity of Streptococci that they can survive for certain periods or replicate in a normal host or host cells such as macrophages. For example, Wiliams and Blakemore (Neuropath. Appl. Neurobiol.: 16, 345-356, 1990; Neuropath. Appl. Neurobiol.: 16, 377-392, 1990; J. Infect. Dis.: 162, 474-481, 1990) show that both polymorphonuclear cells and

macrophage cells are capable of phagocytosing pathogenic S. suis in pigs lacking anti-S. suis antibodies[,]; only pathogenic bacteria could survive and multiply inside macrophages and the pig.

[0053] In a preferred embodiment, the invention, however, provides a deficient or avirulent mutant or vaccine strain which is capable of surviving at least 4-5 days, preferably at least 8-10 days in the host, thereby allowing the development of a solid immune response to subsequent *Streptococcus* infection.

[0054] Due to its persistent but avirulent character, a *Streptococcus* mutant or vaccine strain, as provided by the invention, is well suited to generate specific and/or long-lasting immune responses against Streptococcal antigens. Moreover, possible specific immune responses of the host directed against a capsule are relatively irrelevant because a vaccine strain, as provided by the invention, is typically not recognized by such antibodies.

[0055] In addition, the invention provides a *Streptococcus* vaccine strain according the invention, which strain includes a mutant capable of expressing a *Streptococcus* virulence factor or antigenic determinant.

[0056] In a preferred embodiment, the invention provides a *Streptococcus* vaccine strain, according to the invention, which [strain] includes a mutant capable of expressing a *Streptococcus* virulence factor wherein the virulence factor or antigenic determinant is selected from a group of cellular components, such as muramidase-released protein ([MRP]"MRP"), extracellular factor ([EF]"EF") and cell-membrane associated proteins 60kDA heat shock protein, pneumococcal surface protein A (Psp A), pneumolysin, C protein, protein M, fimbriae, h[a]emagglutinins and [haemolysin]hemolysin or components functionally related thereto.

[9057] In a preferred embodiment, the invention provides a *Streptococcus* vaccine strain [strain comprises] including a mutant capable of over-expressing [said] the virulence factor. In this way, the invention provides a vaccine strain for incorporation in a vaccine which specifically causes a host immune response directed against antigenically important determinants of virulence (listed above), thereby providing specific protection against the determinants. Over-expression can, for example, be achieved by cloning the gene involved behind a strong promoter, which is, for example, constitutionally expressed in a multicopy system, either in a [plsamid]plasmid or via intergration in a genome.

[0058] In yet another embodiment, the invention provides a *Streptococcus* vaccine strain, according to the invention, including a mutant capable of expressing a non-*Streptococcus* protein. Such a vector-*Streptococcus* vaccine strain allows, when used in a vaccine, protection against [other] pathogens other than [Streptococcus] *Streptococcus*.

[0059] Due to its persistent but avirulent character, a *Streptococcus* vaccine strain or mutant as provided by the invention is well suited to generate specific and long-lasting immune responses, not only against Streptococcal antigens, but also against other antigens [when these are] expressed by the strain. Specifically, antigens derived from another pathogen are now expressed without the detrimental effects of the antigen or pathogen which would otherwise have harmed the host.

[0060] An example of such a vector is a *Streptococcus* vaccine strain or mutant wherein the antigen is derived from a pathogen, such as *Actinobacillus pleuropneumonia*, *Mycoplasmatae*, *Bordetella*, *Pasteurella*, *E. coli*, *Salmonella*, *Campylobacter*, *Serpulina* and others.

[0061] The invention also provides a vaccine including a *Streptococcus* vaccine strain or mutant according to the invention and a pharmaceutically acceptable carrier or adjuvant. Carriers or adjuvants are well known in the art[,]; examples are phosphate buffered saline, physiological salt solutions, (double-) [oil-in-water-emulsions]oil-in-water emulsions, aluminumhydroxide, Specol, block- or co-polymers, and others.

[0062] A vaccine according to the invention can include a vaccine strain either in a killed or live form. For example, a killed vaccine including a strain having (over) expressed a Streptococcal or heterologous antigen or virulence factor is very well suited for eliciting an immune response. In a preferred embodiment, the invention provides a vaccine wherein the strain is live, due to its persistent but avirulent character[,]; a Streptococcus vaccine strain, as provided by the invention, is well suited to generate specific and long-lasting immune responses.

[0063] The invention also provides a method for controlling or eradicating a Streptococcal disease in a population comprising vaccinating subjects in the population with a vaccine according to the invention.

[0064] In a preferred embodiment, a method for controlling or eradicating a Streptococcal disease is provided including testing a sample, such as a blood sample, or nasal or

throat swab, f[a]eces, urine, or other samples such as can be sampled at or after slaughter, collected from at least one subject, such as an infant or a pig, in a population partly or wholly vaccinated with a vaccine according to the invention for the presence of encapsulated Streptococcal strains or mutants. Since a vaccine strain or mutant according to the invention is not pathogenic, and can be distinguished from wild-type strains by capsular expression, the detection of (fully) encapsulated Streptococcal strains indicates that wild-type infections are still present. Such wild-type infected subjects can then be isolated from the remainder of the population until the infection has passed. With domestic animals, such as pigs, it is even possible to remove the infected subject from the population as a whole by culling. Detection of wild-type strains can be achieved via traditional culturing techniques, or by rapid detection techniques such as PCR detection.

[0065] In yet another embodiment, the invention provides a method for controlling or eradicating a Streptococcal disease including testing a sample collected from at least one subject in a population partly or wholly vaccinated with a vaccine according to the invention for the presence of capsule-specific antibodies directed against Streptococcal strains. Capsule specific antibodies can be detected with classical techniques known in the art, such as used for Lancefield's group typing or serotyping.

[0066] A preferred embodiment for controlling or eradicating a Streptococcal disease in a population includes vaccinating subjects in the population with a vaccine according to the invention and testing a sample collected from at least one subject in the population for the presence of encapsulated Streptococcal strains and/or for the presence of capsule-specific antibodies directed against Streptococcal strains.

[0067] For example, a method is provided wherein the Streptococcal disease is caused by *Streptococcus suis*.

[0068] The invention also provides a diagnostic assay for testing a sample for use in a method according to the invention [comprising] <u>including</u> at least one means for the detection of encapsulated Streptococcal strains and/or for the detection of capsule-specific antibodies directed against Streptococcal strains.

[0069] The invention further provides a vaccine including an antigen according to the invention and a suitable carrier or adjuvant. The immunogenicity of a capsular antigen provided by the invention is, for example, increased by linking to a carrier (such as a carrier protein), allowing the recruitment of T-cell help in developing an immune response.

[0070] The invention further provides a recombinant micro[-]organism provided with at least a part of a capsular gene cluster derived from Streptococcus suis. The invention provides, for example, a lactic acid bacterium provided with at least a part of a capsular gene cluster derived from Streptococcus suis. Various food-grade lactic acid bacteria (Lactococcus lactis, Lactobacillus casei, Lactobacillus plantarium and Streptococcus gordonii) have been used as delivery systems for mucosal immunization. It has now been shown that oral (or mucosal) administration of recombinant L. lactis, Lactobacillus, and Streptococcus gordonii can elicit local IgA and/or IgG antibody responses to an expressed antigen. The use of oral routes for immunization against infective diseases is desirable because oral vaccines are easier to administer[,] and have higher compliance rates, and because mucosal surfaces are the portals of entry for many pathogenic microbial agents. It is within the skill of the artisan to provide such micro-organisms with (additional) genes.

[0071] The invention further provides a recombinant *Streptococcus suis* mutant provided with a modified capsular gene cluster. It is within the skill of the artisan to swap genes within a Species. In a preferred embodiment, an avirulent *Streptococcus suis* mutant is selected to be provided with at least a part of a modified capsular gene cluster according to the invention.

[0072] The invention further provides a vaccine including a micro[-]organism or a mutant provided by the invention. An advantage of such a vaccine over currently used vaccines is that they include accurately defined micro[-]organisms and well-[characterised]characterized antigens, allowing accurate determination of immune responses against various antigens of choice.

[0073] The invention is further explained in the experimental part of this description without limiting the invention thereto.

# **Description of the Figures**

[0074] FIG. 1 illustrates the organization of the cps2 gene cluster of S. suis type 2.

- (A) Genetic map of the cps2 gene cluster. The shadowed arrows represent potential ORFs. Interrupted ORFs indicate the presence of stop codons or frame-shift mutations. Gene designations are indicated below the ORFs. The closed arrows indicate the position of the potential promoter sequences. I indicates the position of the potential transcription regulator sequence. III indicates the position of the 100-bp repeated sequence.
  - (B) Physical map of the cps2 locus. Restriction sites are as follows: A: AluI; C: ClaI; E[,]: EcoRI; H[,]: HindIII; K[,]: KpnI; M[,]: MluI; N[,]: NsiI; P[.]: PstI; S[.]: SnaBI; Sa: SacI; X[,]: XbaI.
  - (C) The DNA fragments cloned in the various plasmids.
- [0075] FIG. 2 illustrates ethidium bromide stained agarose gel showing PCR products obtained with chromosomal DNA of S. suis strains belonging to the serotypes 1,2, ½, 9 and 14 and cps2J, cpsII, and cps9H primer sets as described herein.
  - (A) cpsII primers; (B) cps2J primers and (C) cps9H primers.
- Lanes 1-3: serotype 1 strains; lanes 4-6: serotype 2 strains; lanes 7-9: serotype 1/2 strains; lanes 10-12: serotype 9 strains and lanes 13-15: serotype 14 strains.
  - (B) Ethidium bromide stained agarose gel showing PCR products obtained with tonsillar swabs collected from pigs carrying S. suis type 2, type 1 or type 9 strains and cps2J, cpsII and cpsH primer sets as described in Materials and Methods. Bacterial DNA suitable for PCR was prepared by using the multiscreen methods as described previously (20).
  - ([A]C) cps[I]II primers. (B) cps2J primers and (C) cps9H primers.

Lanes 1-3: PCR products obtained with tonsillar swabs collected from pigs carrying S. suis type 1 strains; lanes 4-6: PCR [products] products obtained with tonsillar swabs collected from pigs carrying S. suis type 2 strains; lanes 7-9: PCR [products] products obtained with tonsillar swabs collected from pigs carrying S. suis type 9 strains; lanes 10-12: PCR products obtained with chromosomal DNA from serotype 9, 2 and 1 strains respectively; lane 13: negative control, no DNA present.

[0076] FIG. 3 illustrates the CPS2 nucleotide sequences and corresponding amino acid sequences from the open reading frames.

[0077] FIG. 4 illustrates the CPS1 nucleotide sequences and corresponding amino acid sequences from the open reading frames.

[0078] FIG. 5 illustrates the CPS9 nucleotide sequences and corresponding amino acid sequences from the open reading frames.

[0079] FIG. 6 illustrates the CPS7 nucleotide sequences and corresponding amino acid sequences from the open reading frames.

[0080] FIG. 7 illustrates alignment of the N-terminal parts of Cps2J and Cps2K.

Identical amino acids are marked by bars. The amino acids shown in bold are also conserved in CPS14I Cps[l]14J of S. pneumoniae and several other glycosyltransferases (19). The aspartate residues marked by asterisks are strongly conserved.

[0081] FIG. 8 illustrates transmission electron micrographs of thin sections of various S. suis strains.

- (A) wild type strain 10;
- (B) mutant strain l0cpsB;
- (C) mutant strain l0cpsEF.

Bar = 100 nm

[0082] FIG. 9 illustrates the kinetics of phagocytosis of wild type and mutant S. suis strains.

- (A) Kinetics of phagocytosis of wild type and mutant S. suis strains by porcine alveola[i]r macrophages. Phagocytosis was determined as described herein. The Y-axis represents the number of CFU per milliliter in the supernatant fluids as determined by plate counting, the X-axis represents time in minutes.
  - □ wild type strain 10;
  - o mutant strain 10cpsB;
  - Δ mutant strain l0cpsEF.
- (B) Kinetics of intracellular killing of wild type and mutant S. suis strains by porcine AM. The intracellular killing was determined as described herein. The Y-axis represents the

number of CFU per ml in the supernatant fluids after lysis of the macrophages as determined by plate counting, the X-axis represents time in minutes.

- □ wild type strain 10;
- o mutant strain 10cpsB;
- Δ mutant strain l0cpsEF.

[0083] FIG. 10 illustrates the nucleotide sequence alignment of the highly conserved 100-bp repeated element.

- 1) 100-bp repeat between cps2G and cps2H
- 2) 100[—]-bp repeat within "cps2M"
- 3) 100[—]-bp repeat between cps2O and cps2P

[0084] FIG. 11 illustrates the cps2, cps9 and cps7 gene clusters of S. suis serotypes 2, 9 and 7.

- (A) Genetic organization of the cps2 gene cluster [84]. The large arrows represent potential ORFs. Gene designations are indicated below the ORFs. Identically filled arrows represent ORFs which showed homology. The small closed arrows indicate the position of the potential promoter sequences. I indicates the position of the potential transcription regulator sequence.
- (B) Physical map and genetic organization of the cps9 gene cluster [15]. Restriction sites are as follows: B: BamHI; P: PstI; H: HindIII; X: XbaI. The DNA fragments cloned in the various plasmids are indicated. The open arrows represent potential [OREs] ORFs.
- (C) Physical map and genetic organization of the [cps7gene]cps7 gene cluster. Restriction sites are as follows: C: Clal; P: PstI; Sc: ScaI. The DNA fragments cloned in the various plasmids are indicated. The open arrows represent potential OR[E]Fs.

[0085] FIG. 12 illustrates [Ethidium] ethidium bromide stained agarose gel showing PCR products.

- (A) Ethidium bromide stained agarose gel showing PCR products obtained with chromosomal DNA of S. suis strains belonging to the serotypes 1, 2, 9 and 7 and the cps7H primer set. Strain designations are indicated above the lanes. C: negative control, no DNA present. M: molecular size marker (lambda digested with *EcoRI* and *HindIII*).
  - (B) Ethidium bromide stained agarose gel showing PCR products obtained with serotype 7 strains collected in different countries and from different organs. Bacterial DNA suitable for PCR was prepared by using the multiscreen method as described herein [89]. Strain designations are indicated above the lanes. M: molecular size marker (lambda digested with *EcoRI* and HindIII).

## **Detailed Description of the Invention**

## Experimental part

#### MATERIAL AND METHODS

#### Bacterial strains and growth conditions.

[0086] The bacterial strains and plasmids used in this study are listed in Table 1. S. suis strains were grown in Todd-Hewitt broth (code CM189, Oxoid), and plated on Columbia agar blood base (code CM331, Oxoid) containing 6% (v/v) horse blood. E. coli strains were grown in Luria broth (28) and plated on Luria broth containing 1.5% (w/v) agar. If required, antibiotics were added to the plates at the following concentrations: spectinomycin: 100 ug/ml for S. suis and 50 ug/ml for E. coli and ampicillin, 50 ug/ml.

[0087] Serotyping. The S. suis Strains were [serotypes] serotyped by the slide agglutination test with serotype-specific antibodies (44).

[0088] DNA techniques. Routine DNA manipulations were performed as described by Sambrook []et al. (36).

[0089] Alkaline phosphatase activity. To screen for PhoA fusions in *E. coli*, plasmid libraries were constructed. Therefore, chromosomal DNA of *S. suis* type 2 was digested with *Alul*. The 300-500-bp fragments were ligated to SmaI-digested pPHOS2. Ligation mixtures were transformed to the PhoA<sup>-</sup> *E. coli* strain CC118. Transformants were plated on LB media

supplemented with 5-Bromo-4-chloro-3-indolylfosfaat (BCIP, 50 ug/ml, Boehringer, Mannheim, Germany). Blue colonies were purified on fresh LB/BCIP plates to verify the blue phenotype.

[0090] DNA sequence analysis. DNA sequences were determined on a 373A DNA Sequencing System (Applied Biosystems, Warrington, GB). Samples were prepared by using an ABI/PRISM dye terminator cycle sequencing ready reaction kit (Applied Biosystems). Sequencing data were assembled and analyzed using the MacMollyTetra program. Custom-made sequencing primers were purchased from Life Technologies. Hydrophobic stretches within proteins were predicted by the method of Klein []et al. (17). The BLAST program available on Netscape Navigator<sup>TM</sup> was used to search for protein sequences related to the deduced amino acid sequences.

[0091] Construction of gene-specific knock-out mutants of *S. suis*. To construct the mutant strains 10cpsB and 10cpsEF, we electrotransformed the pathogenic serotype 2 strain 10 (45, 49) of *S. suis* with pCPS11 and pCPS28 respectively. In these plasmids, the *cpsB* and cpsEF genes were disturbed by the insertion of a spectinomycin-resistance gene. To create pCPS11, the internal 400 bp *PstIBamHI* fragment of the *cpsB* gene in pCPS7 was replaced by the Spc<sup>R</sup> gene. For this purpose, pCPS7 was digested with *PstI* and BamHI and ligated to the 1,200-bp PstI-*BamHI* fragment, containing the Spc<sup>R</sup> gene, from pIC-spc. To construct pCPS28, we have used pIC20R. In this plasmid we inserted the *KpnI-SalI* fragment from pCPS17 (resulting in pCPS25) and the *XbaI-ClaI* fragment from pCPS20 (resulting in pCPS27). pCPS27 was digested with *PstI* and *XhoI* and ligated to the 1,200-bp *PstI-XhoI* fragment, containing the Spc<sup>R</sup> gene of pIC-spc. The electrotransformation to *S. suis* was carried out as described before (38).

[0092] Southern blotting and hybridization. Chromosomal DNA was isolated as described by Sambrook []et al. (36). DNA fragments were separated on 0.8% agarose gels and transferred to Zeta-Probe GT membranes (Bio-Rad) as described by Sambrook et al. (36). DNA probes were [labelled] labeled with [( -32P] dCTP (3000 Ci mmol<sup>-1</sup>; Amersham) by use of a random primed labeling kit (Boehringer). The DNA on the blots was hybridized at 65°C with appropriate DNA probes as recommended by the supplier of the Zeta-Probe membranes. After hybridization, the membranes were washed twice with a solution of 40 mM sodium phosphate, pH 7.2, 1 mM

EDTA, 5% SDS for 30 min at 65°C and twice with a solution of 40 mM sodium phosphate, pH 7.2, 1 mM EDTA, 1% SDS for 30 min at 65°C.

PCR. The primers used in the cps2J PCR correspond to the positions 13791-13813 and 14465-14443 in the S. suis cps2 locus. The sequences were: CAAACGCAAGGAATTACGGTATC-3' (SEQ. 5'-ID. No. 1) and GAGTATCTAAAGAATGCCTATTG-3' (SEQ. ID. No. 2). The primers used for the cpslI PCR correspond to the positions 4398-4417 and 4839-4821 in the S. suis cps1 sequence. The sequences were: 5'-GGCGGTCTAGCAGATGCTCG-3' (SEQ. ID. No. 3) and 5' -GCGAACTGTTAGCAATGAC-3' (SEQ. ID. No. 4). The primers used in the cps9H PCR correspond to the positions 4406-4126 and 4494-4475 in the S. suis cps9 sequence. The sequences 5'-GGCTACATATAATGGAAGCCC3' (SEQ. ID No. CGGAAGTATCTGGGCTACTG-3' (SEQ. ID. No. 6).

[0094] Construction of gene-specific knock-out mutants of *S. suis*. To construct the mutant strains 10cpsB and 10cpsEF, we electrotransformed the pathogenic serotype 2 strain 10 of *S. suis* with pCPS11 and pCPS28 respectively. In these plasmids, the *cpsB* and cpsEF genes were disturbed by the insertion of a spectinomycin-resistance gene. To create pCPS11, the internal 400 bp *PstI-BamHI* fragment of the *cpsB* gene in pCPS7 was replaced by the Spc<sup>R</sup> gene. For this purpose, pCPS7 was digested with *PstI* and BamHI and ligated to the 1,200-bp *PstI-BamHI* fragment, containing the Spc<sup>R</sup> gene, from pIC-spc. To construct pCPS28, we have used pIC20R. In this plasmid, we inserted the *KpnI-SalI* fragment from pCPS17 (resulting in pCPS25) and the *XbaI-ClaI* fragment from pCPS20 (resulting in pCPS27). pCPS27 was digested with *PstI* and *XhoI* and ligated to the 1,200-bp *PstI-XhoI* fragment, containing the Spc<sup>R</sup> gene of pIC-spc. The electrotransformation to *S. suis* was carried out as described before (38).

[]et al. (23). Briefly, to opsonize the cells,  $10^7$  S. suis cells were incubated with 6% SPF-pig serum for 30 min at  $37^{\circ}$ C in a head-over-head rotor at 6 rpm.  $10^7$  AM and  $10^7$  opsonized S. suis cells were combined and incubated at  $37^{\circ}$ C under continuous rotation at 6 rpm. At 0, 30, 60 and 90 min, 1- ml samples were collected and mixed with 4 ml of ice-cold EMEM to stop phagocytosis. Phagocytes were removed by centrifugation for 4 min at  $110 \times g$  and  $4^{\circ}$ C. The number of colony-

forming units, ([CFU]"CFU") in the supernatants was determined. Control experiments were carried out simultaneously by combining 10<sup>7</sup> opsonized S. suis cells with EMEM (without AM).

[0096] Killing assays. AM (10<sup>7</sup>/ml) and opsonized *S. suis* cells (10<sup>7</sup>/ml) were mixed 1:1 and incubated for 10 min at 37°C under continuous rotation at 6 rpm. Ice-cold EMEM was added to stop further phagocytosis and killing. To remove extracellular *S. suis* cells, phagocytes were washed twice (4 min, 110 x g, 4°C) and resuspended in 5 ml EMEM containing 6% SPF serum. The tubes were incubated at 37°C under rotation at 6 rpm. After 0, 15, 30, 60 and 90 min, samples were collected and mixed with ice-cold EMEM to stop further killing. The samples were centrifuged for 4 min at 110 x g at 4°C and the phagocytic cells were lysed in EMEM containing 1% saponine for 20 min at room temperature. The number of CFU in the suspensions was determined.

[0097] Pigs. Germfree pigs, cross[-]breeds of Great Yorkshire and Dutch [l]Landrace, were obtained from sows by caesarian sections. The surgery was performed in sterile flexible film isolators. Pigs were allotted to groups, each consisting of 4 pigs, and were housed in sterile stainless steel incubators.

as described before. To predispose the pigs for infection with *S. suis*, five-day old pigs were inoculated intranasally with about 10<sup>7</sup> CFU of *Bordetella bronchiseptica* strain 92932. Two days later, the pigs were inoculated intranasally with *S. suis* type 2 (10<sup>6</sup> CFU). Pigs were monitored twice daily for clinical signs of disease, such as fever, nervous signs and lameness. Blood samples were collected three times a week from each pig. White blood cells were counted with a cell counter. To monitor infection with *S. suis* and *B. bronchiseptica* and to check for absence of contaminants, we collected swabs of nasopharynx and feces daily. The swabs were plated directly onto Columbia agar containing 6% horse blood. After three weeks, the pigs were killed and examined for pathological changes. Tissue specimens from the central nervous system, serosae, and joints were examined bacteriologically and histologically as described herein (45, 49). Colonization of the serosae was scored positively when *S. suis* was isolated from the pericardium, thoracal pleura or the peritoneum. Colonization of the joints was scored positively when *S. suis* was isolated from one or more joints (12 joints per animal were scored).

[0099] Vaccination and challenge. One week old pigs were vaccinated intravenously with a dosage of 106 cfu of the *S. suis* strains 10cpsEF or 10cpsB. Three weeks later, the pigs were challenged intravenously with the pathogenic Serotype 2 strain 10 (107 cfu). Disease monitoring, [haematologicl]hematological, serological and bacteriological examinations as well as post-mortum examinations were as described before under experimental infections.

[0100] Electron Microscopy. Bacteria were prepared for electron microscopy as described by Wagenaar []et al. (50). Shortly, bacteria were mixed with agarose MP (Boehringer) of 37° C to a concentration of 0.7%. The mixture was immediately cooled on ice. Upon gelifying, samples were cut into 1 to 1.5 mm slices and incubated in a fixative containing 0.8% glutaraldehyde and 0.8% osmiumtetraoxide. Subsequently, the samples were fixed and stained with uranyl acetate by microwave stimulation, dehydrated and imbedded in eponaraldite resin. Ultra-thin sections were counterstained with lead citrate and examined with a Philips CM 10 electron microscope at 80 kV (FIG. 8).

[0101] Isolation of porcine alveolar macrophages (AM). Porcine AM were obtained from the lungs of specific pathogen free ([SPF]"SPF") pigs. Lung lavage samples were collected as described by van Leengoed et al. (43). Cells were suspended in EMEM containing 6% (v/v). SPF-pig serum and adjusted to 10<sup>7</sup> cells per ml.

#### RESULTS

#### Identification of the cps locus.

[0102] The cps locus of S. suis type 2 was identified through [by making use of] a strategy developed for the genetic identification of exported proteins (13, 31). In this system, we used a plasmid (pPHOS2) containing a truncated alkaline phosphatase gene (13). The gene lacked the promoter sequence, the translational start site and the signal sequence. The truncated gene is preceded by a unique SmaI restriction site. Chromosomal DNA of S. suis type 2, digested with AluI, was randomly cloned in this restriction site. Because translocation of PhoA across the cytoplasmic membrane of E. coli is required for enzymatic activity, the system can be used to select for S. suis fragments containing a promoter sequence, a translational start site and a functional signal sequence. Among 560 individual E. coli clones tested, 16 displayed a dark blue

phenotype when plated on media containing BCIP. DNA sequence analysis of the inserts from several of these plasmids [were] was performed (results not shown) and the deduced amino acid sequences were analyzed. The hydrophobicity profile of one of the clones (pPHOS7, results not shown) showed that the N-terminal part of the sequence resembled the characteristics of a typical signal peptide: a short hydrophilic N-terminal region is followed by a hydrophobic region of 38 amino acids. These data indicate that the phoA system was successfully used for the selection of S. suis genes encoding exported proteins. Moreover, the sequences were analyzed for similarities present in the databases. The sequence of pPHOS7 showed a high similarity (37% identity) with the protein encoded by the cps14C gene of Streptococcus pneumoniae (19). This strongly suggests that pPHOS7 contains a part of the cps operon of S. suis type 2.

[0103] Cloning of the flanking cps genes. In order to clone the flanking cps genes of S. suis type 2, the insert of pPHOS7 was used as a probe to identify chromosomal DNA fragments which contain flanking cps genes. A 6-kb HindIII fragment was identified and cloned in pKUN19. This yielded clone pCPS6 (FIG. 1, part C). Sequence analysis of the insert of pCPS6 revealed that pCPS6 most probably contained the 5'-end of the cps locus, but still lacked the 3'-end. Therefore, sequences of the 3'-end of pCPS6 were in turn used as a probe to identify chromosomal fragments containing cps sequences located further downstream. These fragments were also cloned in pKUN19, resulting in pCPS17. Using the same system of chromosomal walking, we subsequently generated the plasmids pCPS18, pCPS20, pCPS23 and pCPS26, containing downstream cps sequences.

[0104] Analysis of the cps operon. The complete nucleotide sequence of the cloned fragments was determined (FIG. 4). Examination of the compiled sequence revealed the presence of at least 13 potential open reading frames (Orfs), which were designated as Orf 2Y, Orf2X and Cps2A-Cps2K (FIG. 1, part A; FIG. 1, part A). Moreover, a 14th, incomplete[,] Orf (Orf 2Z) was located at the 5'-end of the sequence. Two potential promoter sequences were identified. One was located 313 bp (locations 1885-1865 and 1884-1889) upstream of Orf2X. The other potential promoter sequence was located 68 bp upstream of Orf2Y (locations 2241-2236 and 2216-2211). Orf2Y is expressed in opposite orientation. Between Orfs 2Y and 2Z, the sequence contained a potential stem-loop structure, which could act as a transcription terminator. Each Orf is preceded

by a ribosome-binding site and the majority of the Orfs are very closely linked. The only significant intergenic gap was found between Cps2G and Cps2H (389 nucleotides). However, no obvious promoter sequences or potential stem-loop structures were found in this region. These data suggest that Orf2X and Cps2A-Cps2K are arranged as an operon.

[0105] An overview of all Orfs with their properties is shown in Table 2. The majority of the predicted gene products is related to proteins involved in polysaccharide biosynthesis. Orf2Z showed some similarity with the YitS protein of *Bacillus subtilis*. YitS was identified during the sequence analysis of the complete genome of B. *subtilis*. The function of the protein is unknown.

[0106] Orf2Y showed similarity with the YcxD protein of B. subtilis (53). Based on the similarity between YcxD and MocR of Rhizobium meliloti (33), YcxD was suggested to be a regulatory protein.

[0107] Orf2X showed similarity with the hypothetical YAAA proteins of *Haemophilus* influenzae and E. coli. The function of these proteins is unknown.

[0108] The gene products encoded by the cps2A, cps2B, cps2C and cps2D genes showed approximate similarity with the CpsA, CpsC, CpsD and CpsB proteins of several serotypes of Streptococcus pneumoniae (19), respectively. This suggests similar functions for these proteins. Hence, Cps2A may have a role in the regulation of the capsular polysaccharide synthesis. Cps2B and Cps2C could be involved in the chain length determination of the type 2 capsule and Cps2C can play an additional role in the export of the polysaccharide. The Cps2D protein of S. suis is related to the CpsB protein of S. pneumoniae and to proteins encoded by genes of several other Gram-positive bacteria involved in polysaccharide or exopolysaccharide synthesis, but their function is unknown (19).

[0109] The protein encoded by the cps2E gene showed similarity to several bacterial proteins with [glycosyl transferase]glycosyltransferase activities Cps14E and Cps19fE of S. pneumoniae serotypes 14 and 19F (18, 19, 29), CpsE of Streptococcus salvarius (X94980) and CpsD of Streptococcus agalactiae (34). Recently, Kolkman et al. (18) showed that Cps14E is a glucosyl-1-phosphate transferase that links glucose to a lipid carrier, the first step in the

biosynthesis of the S. pneumoniae type 14 repeating unit. Based on these data, a similar function may be fulfilled by Cps2E of S. suis.

[0110] The protein encoded by the cps2F gene showed similarity to the protein encoded by the rfbU gene of Salmonella enteritica. (25). This similarity is most pronounced in the C-terminal regions of these proteins. The rfbU gene was shown to encode[d] mannosyltransferase activity (25).

[0111] The cps2G gene encoded a protein that showed moderate similarity with the rfbF gene product of Campylobacter hyoilei (22), the epsF gene product of S. thermophilus (40) and the capM gene product of S. aureus (24). On the basis of similarity, the rfbF, epsF and capM genes are suggested to encode[d] galactosyltransferase activities. Hence, a similar [glycosyltransferase]glycosyltransferase activity could be fulfilled by the cps2G gene product.

[0112] The cps2H gene encodes a protein that is similar to the N-terminal region of the lgtD gene product of Haemophilus influenzae (U32768). Moreover, the hydrophobicity plots of Cps2H and LgtD looked very similar in these regions (data not shown). Based on sequence similarity, the lgtD gene product was suggested to have [glycosyl transferase]glycosyltransferase activity (U32768).

[0113] The gene product encoded by the *cps2I* gene showed some similarity with a protein of *Actinobacillus actinomycetemcomitans* (AB002668). This protein is part of the gene cluster responsible for the serotype-b-specific antigen of *A. actinomycetemcomitans*. The function of the protein is unknown.

[0114] The gene products encoded by the *cps2J* and *cps2K* genes showed significant similarities to the Cpsl4J protein of *S. pneumoniae*. The *cps14J* gene of *S. pneumoniae* was shown to encode a β-1,4-galactosyltransferase activity. In *S. pneumoniae*. CpsJ is responsible for the addition of the fourth (*i.e.* last) sugar in the synthesis of the *S. pneumoniae* serotype 14 polysaccharide (20). Even some similarity was found between Cps2J and Cps2K (FIG. 2, 25.5% similarity). This similarity was most pronounced in the N-terminal regions of the proteins (FIG. 7). Recently, two small conserved regions were identified in the N-terminus of Cps14J and Cps14I and their homologues (20). These regions were predicted to be important for catalytic activity. Both regions, DXS and DXDD [Fig.] (FIG. 2), were also found in Cps2J and Cps2K.

[0115]Distribution of the cps2 genes in other S. suiss serotypes. To examine the relationship between the cps2 genes and cps genes in the other S. suis serotypes, we performed crosshybridization experiments. DNA fragments of the individual cps2 genes were amplified by PCR, labeled with <sup>32</sup>P, and used to probe Southern blots of chromosomal DNA of the reference strains of the 35 different S. suis serotypes. Large variations in the hybridization patterns were observed (Table 4). As a positive control, we used a probe specific for 16S rRNA. The 16S rRNA probe hybridized with all serotypes tested. However, none of the other genes tested were common in all serotypes. Based on the genetic organization of the genes, we previously suggested that orfX and cpsA-cpsK genes are part of one operon and that the proteins encoded by these genes are all involved in polysaccharide biosynthesis. OrfY and OrfZ are not a part of this operon, and their role in the polysaccharide biosynthesis is unclear. Based on sequence similarity data, OrfY may be involved in regulation of the cps2 genes. OrfZ is proposed to be unrelated to polysaccharide biosynthesis. Probes specific for the orfZ, orfY, orfX, cpsA, cpsB, cpsC and cpsD genes hybridized with most other serotypes. This suggests that the proteing encoded by these genes are not typespecific, but may perform more common functions in biosynthesis of the capsular polysaccharide. This confirms previous data which showed that the cps2A-cps2D genes showed strong similarity to cps genes of several serotypes of Streptococcus pneumoniae. Based on this similarity, Cps2A is possibly a regulatory protein, whereas Cps2B and Cps2C may play a role in length determination and export of polysaccharide. The cps2E gene hybridized with DNA of Serotypes 1, 2, 14 and 1/2. The cps2E gene showed a strong similarity to the cps14E gene of S. pneumoniae (18). This enzyme was shown to have a glucosyl-1-phosphate activity and catalyzed the transfer of glucose to a lipid carrier (18). These data indicate that a glycosyltransferase closely related to Cps14E may be responsible for the first step in the biosynthesis of polysaccharide in the S. suis serotypes 1, 2, 14 and 1/2. The cps2F, cps2G, cps2H, cps2I and cps2J genes hybridized with chromosomal DNA of serotypes 2 and 1/2 only. The cps2G gene showed an additional weak hybridization signal with DNA of serotype 34. In agglutination tests, serotype 1/2 showed agglutination with sera specific for serotype 2 as well as with sera specific for serotype 1. This suggests that serotype 1/2 shares antigenic determinants with both types 1 and 2. The hybridization data confirmed these data. All putative glycosyltransferases present in serotype 2 are also present in serotype 1/2. The cps2K

gene showed a [similar ]hybridization pattern [as]similar to the cps2E gene. Hybridization was observed with DNA of serotypes 1, 2, 14 and 1/2. Taken together, these hybridization data show that the cps2 gene cluster can be divided into three regions: a central region containing the type-specific genes is flanked by two regions containing common genes for various serotypes.

[0116] Cloning of the type-specific cps genes of serotypes 1 and 9. To clone the type-specific cps genes of S. suis serotype 1, we used the cps2E gene as a probe to identify chromosomal DNA fragments of type 1 which contain flanking cps genes. A 5 kb EcoRV fragment was identified and cloned in pKUN19. This yielded pCPS1-1 (FIG. 1, part B). This fragment was in turn used as a probe to identify an overlapping 2.2 kb HindIII fragment. pKUN19 containing this HindIII fragment was designated pCPS1-2. The same strategy was followed to identify and clone the type-specific cps genes of serotype 9. In this case, we used the cps2D gene as a probe. A 0.8 kb HindIII-XbaI fragment was identified and cloned, yielding pCPS9-1 (FIG. 1, part C). This fragment was in turn used as a probe to identify a 4 kb XbaI fragment. pKUN19 containing this 4 kb XbaI fragment was designated pCPS9-2.

[0117] Analysis of the cloned cps1 genes. The complete nucleotide sequence of the inserts of pCPS1-1 and pCPS1-2 was determined (FIG. 5). Examination of the sequence revealed the presence of five complete and two incomplete Orfs (FIG.1, part B). Each Orf is preceded by a ribosome-binding site. In accord with data obtained for the cps2 genes of serotype 2, the majority of the Orfs is very closely linked. The only significant gap (718 bp) was found between Cps1G and Cps1H. No obvious promoter sequences or potential stem-loop structures could be found in this region. This suggests that, as in serotype 2, the cps genes in serotype 1 are arranged in an operon.

[0118] An overview of the Orfs and their properties [in] is shown in Table 2. As expected on the basis of the hybridization data (Table 4), the protein encoded by the *cps1E* gene was related to Cps2E of *S. suis* type 2 (identity of 86%). The fragment cloned in pCPS1-1 lacked the coding region for the first 7 amino acids of the *cps1E* gene.

[0119] The protein encoded by the *cps1F* and *cps1G* genes showed strong similarity to the Cps14F and Cps14G proteins of *Streptococcus pneumoniae* serotype 14, respectively (20). The function of the Cps14F is not completely clear, but it has been suggested that Cps14F [can

enhance] has a role in glycosyltransferase activity. The cps14G gene of S. pneumoniae was shown to encode  $\beta$ -1, 4-galactosyltransferase activity. In S. pneumoniae type 14, this activity is required for the second step in the biosynthesis of the oligosaccharide subunit (20). Based on the similarity of the data, similar glycosyltransferase and enhancing activities are suggested for the [cps 1G]cps1G and cps1F genes of S. suis type 1.

[0120] The protein encoded by the *cps1H* gene showed similarity to the Cps14M protein of *S. pneumoniae* (20). Based on sequence similarity, Cps14H was proposed to be the polysaccharide polymerase (20).

[0121] The protein encoded by the cpsII gene showed some similarity with the Cps14J protein of S. pneumoniae (19). The cpsI4J gene was shown to encode a  $\beta$ -1, 4-galactosyltransferase activity, responsible for the addition of the fourth (i.e. last) sugar in the synthesis of the S. pneumoniae serotype 14 polysaccharide.

[0122] Between Cps1G and Cps1H<sub>2</sub> a gap of 718 bp was found. This region revealed three small Orfs. The three Orfs were expressed in three different reading frames and were not preceded by potential ribosome binding sites, nor contained potential start sites. However, the three potential gene products encoded by this region showed some similarity with three successive regions of the C-terminal part of the EpsK protein of *Streptococcus thermophilus* (27% identity, 40). The region related to the first 82 amino acids is lacking.

[0123] Analysis of the cloned cps9 genes. We also determined the complete nucleotide sequence of the inserts of pCPS9-1 and pCPS9-2 (FIG. 6). Examination of the sequence revealed the presence of three complete and two incomplete Orfs (FIG.1, part C). As in serotypes 1 and 2, all Orfs are preceded by a ribosome-binding site and are very closely coupled. As suggested by the hybridization data (Table 4), the Cps2D and Cps9D proteins were highly related (Table 2). Based on sequence comparisons, pCPS9-1 lacked the first 27 amino acids of the Cps9D protein.

[0124] The protein encoded by the *cps9E* gene showed some similarity with the CapD protein of *Staphylococcus aureus* serotype 1 (24). Based on sequence similarity data, the Cap1D protein was suggested to be an epimerase or a dehydratase involved in the synthesis of N-acetylfructosamine or N-acetylgalactosamine (63).

- [0125] Cps9F showed some similarity to the CapM proteins of S. aureus serotypes 5 and 8 (61, 64, 65). Based on sequence similarity data, Cap5M and Cap8M are proposed to be glycosyltransferases (63).
- [0126] The protein encoded by the *cps9G* gene showed some similarity [with ] to a protein of *Actinobacillus actinomycetemcomitans* (AB002668\_4). This protein is part of a gene cluster responsible for the serotype-b specific antigens of *Actinobacillus actinomycetemcomitans*. The function of the protein is unknown.
- [0127] The protein encoded by the *cps9H* gene showed some similarity [with] <u>to</u> the *rfbB* gene of *Yersinia enterolitica* (68). The RfbB protein was shown to be essential for O-antigen synthesis, but the function of the protein in the synthesis of the 0:3 lipopolysaccharide is unknown.
- [0128] Serotype 1 and serotype 9 specific cps genes. To determine whether the cloned fragments in pCPS1-1, pCPS1-2, pCPS9-1 and pCPS9-2 contained the type-specific genes for serotype 1 and 9, respectively, cross[]-hybridization experiments were performed. DNA fragments of the individual cps1 and cps9 genes were amplified by PCR, labeled with <sup>32</sup>P, and used to probe Southern blots of chromosomal DNA of the reference strains of the 35 different S. suis serotypes. The results are shown in Table 5. Based on the data obtained with the cps2E probe (Table 4), the cps1E probe was expected to hybridize with chromosomal DNA of S. suis serotypes 1, 2, 14, 27 and 1/2. The cps1H, cps9E and cps9F probes hybridized with most other serotypes. However, the cps1F and cps1G and cps1I probes hybridized with chromosomal DNA of serotypes 1 and 14 only. The cps9G and cps9H probes hybridized with serotype 9 only. These data suggest that the cps9G and cps9H probes are specific for serotype 9 and, therefore, could be useful tools for the development of rapid and sensitive diagnostic tests for S. suis type 9 infections.
- [0129] Type specific PCR. So far, the probes were tested on the 35 different reference strains only. To test the diagnostic value of the typespecific *CpS* probes further, several other *S. suis* serotype 1, 2, 1/2, 9 and 14 strains were used. Moreover, since a PCR[]-based method would be even more rapid and sensitive than a hybridization test, we tested whether we could use a PCR for the serotyping of the *S. suis* strains. The oligonucleotide primer sets were chosen within the *cps2J*, *cps1I* and *cps9H* genes. Amplified fragments of 675 bp, 380 bp and 390 bp were expected, respectively. The results show that 675 bp fragments were amplified on type 2 and 1/2

strains using *cps2J* primers; 380 bp fragments were amplified on type 1 and 14 strains using *cps1I* primers and 390 bp fragments were amplified on type 9 strains using *cps9H* primers.

[0130] Construction of mutants impaired in capsule production. To evaluate the role of the capsule of S. suis type 2 in the pathogenesis, we constructed two isogenic mutants in which capsule production was disturbed. To construct mutant 10cpsB, pCPS11 was used. In this plasmid, a part of the cps2B gene was replaced by the spectinomycin-resistance gene. To construct mutant strain 10cpsEF, the plasmid pCPS28 was used. In pCPS28, the 3'-end of cps2E gene, as well as the 5'-end, of cps2F gene, were replaced by the spectinomycin-resistance gene. pCPS11 and pCPS28 were used to electrotransform strain 10 of S. suis type 2 and spectinomycinresistant colonies were selected. Southern blotting and hybridization experiments were used to select double [cross over]crossover integration events (results not shown). To test whether the capsular structure of the strains 10cpsB and 10cpsEF was disturbed, we used a slide agglutination test using a suspension of the mutant strains in hyperimmune anti-[S. suis] <u>S. suis</u> type 2 serum (44). The results showed that even in the absence of serotype specific antisera, the bacteria agglutinated. This indicates that, in the mutant strains, the capsular structure was disturbed. To confirm this, thin sections of wild type and mutant strains were compared by electron microscopy. The results showed that, compared to the wild type (FIG. 3, part A), the amount of capsule produced by the mutant strains was greatly reduced (FIG. 3, part B and part C). Almost no capsular material could be detected on the surface of the mutant strains.

[0131] Capsular mutants are sensitive to phagocytosis and killing by porcine alveolar macrophages ("PAM"). The capsular mutants were tested for their ability to resist phagocytosis by PAM in the presence of porcine SPF serum. The wild type strain 10 seemed to be resistant to phagocytosis under these conditions (FIGs. 9A and 9B). In contrast, the mutant strains were efficiently ingested by macrophages (FIGs. 9A and 9B). After 90 min., more than 99.7% (strain 10cpsB) and 99.8% (strain 10cpsEF) of the mutant cells were ingested by the macrophages. Moreover, as shown in FIGs. 9A and 9B the ingested strains were efficiently killed by the macrophages. 90-98% of all ingested cells were killed within 90 min. No differences could be observed between wild type and mutant strains. These data indicate that the capsule of *S. suis* type 2 efficiently protects the organism from uptake by macrophages *in vitro*.

[0132] Capsular mutants are less virulent for germfree piglets. The virulence properties of the wild-type and mutant strains were tested after experimental infection of newborn germfree pigs (45, 49). Table 1 shows that specific and nonspecific signs of disease could be observed in all pigs inoculated with the wild type strain. Moreover, all pigs inoculated with the wild type strain died during the course of the experiment or were killed because of serious illness or nervous disorders (Table 3). In contrast, the pigs inoculated with strains 10cpsB and 10cpsEF showed no specific signs of disease and all pigs survived until the end of the experiment (Table 6). The temperature of the pigs inoculated with the wild type strain increased 2 days after inoculation and remained high until day 5 (Table 3). The temperature of the pigs inoculated with the mutant strains sometimes exceeded the 40°C, however, we could observe significant differences in the fever index (i.e. percent of observations in an experimental group during which pigs showed fever (>40°C)) between pigs inoculated with wild type and mutant strains. All pigs showed increased numbers of polymorphonuclear leucocytes (PMLs) (>10 x 109 PMLs per litre) (Table 3). However, in pigs inoculated with the mutant strains, the percentage of samples with increased numbers of PMLs was considerably lower. S. suis strains and B. bronchiseptica could be isolated from the nasopharynx and feces swab samples of all pigs from 1 day post-infection until the end of the experiment (Table 3). Postmortem, the wild type strain could frequently be isolated from the central nervous system ([CNS]"CNS"), kidney, heart, liver, spleen, serosae, joints and tonsils. Mutant strains could easily be recovered [form] from the tonsils, but were never recovered from the kidney, liver or spleen. Interestingly, low numbers of the mutant strains were isolated from the CNS, the serosae, the joints, the lungs and the heart. Taken together, these data strongly indicated that mutant S. suis strains, impaired in capsule production, are not virulent for young germfree pigs.

[0133] We describe the identification and the molecular characterization of the *cps* locus, involved in the capsular polysaccharide biosynthesis, of *S. suis*. Most of the genes seemed to belong to a single transcriptional unit, suggesting a co[-]ordinate control of these genes. We assigned functions to most of the gene products. We thereby identified regions involved in regulation (Cps2A), chain length determination (Cps2B, C), export (Cps2C) and biosynthesis (Cps2E, F, G, H, J, K). The region involved in biosynthesis is located at the center of the gene

cluster and is flanked by two regions containing genes with more common functions. The incomplete or f2Z gene was located at the 5'-end of the cloned fragment. Or f2Z showed some similarity with the YitS protein of B. subtilis. However, because the function of the YitS protein is unknown, this did not give us any information about the possible function of Or f2Z. Because the or f2Z gene is not a part of the cps operon, a role of this gene in polysaccharide biosynthesis is not expected. The Or f2Y protein showed some similarity with the YcxD protein of B. subtilis (53). The YcxD protein was suggested to be a regulatory protein. Similarly, Or f2Y may be involved in the regulation of polysaccharide biosynthesis. The Or f2X protein showed similarity with the YAAA proteins of H. influenzae and E. coli. The function of these proteins is unknown. In S. [suis] suis type 2, the or f2X gene seemed to be the first gene in the cps2 operon. This suggests a role of Or f2X in the polysaccharide biosynthesis. In H. influenzae and E. coli, however, these proteins are not associated with capsular gene clusters. The analysis of isogenic mutants impaired in the expression of Or f2X should give more insight in the presumed role of Or f2X in the polysaccharide biosynthesis of S. suis type 2.

[0134] The gene products encoded by the cps2E, cps2F, cps2G, cps2H, cps2J and cps2K genes showed little similarity with glycosyltransferases of several Gram-positive or Gramnegative bacteria (18, 19, 20, 22, 25). The cps2E gene product shows some similarity with the Cps14E protein of S. pneumoniae (18, 19). Cps14E is a glucosyl-1-phosphate transferase that links glucose to a lipid carrier (18). In S. pneumoniae, this is the first step in the biosynthesis of the oligosaccharide repeating unit. The structure of the S. suis serotype 2 capsule contains glucose, galactose, rhamnose, N-acetyl glucos[e]amine and sialic acid in a ratio of 3:1:1:1:1 (7). Based on these data, we conclude that Cps2E of S. suis has glucosyltransferase activity[,] and is involved in the linkage of the first sugar to the lipid carrier.

[0135] The C-terminal region of the *cps2F* gene product showed some similarity with the RfbU of *Salmonella enteritica*. RfbU was shown to have mannosyltransferase activity (24). Because mannosyl is not a component of the *S. suis* type 2[,] polysaccharide, a mannosyltransferase activity is not expected in this organism. Nevertheless, *cps2F* encodes a glycosyltransferase with another sugar specificity.

[0136] Cps2G showed moderate similarity to a family of gene products suggested to encode galactosyltransferase activities (22, 24, 40). Hence, a similar activity is shown for Cps2G.

[0137] Cps2H showed some similarity with LgtD of *H. influenzae* (U32768). Because LgtD was proposed to have glycosyltransferase activity[], a similar activity is fulfilled by Cps2H.

[0138] Cps2J and Cps2K showed similarity to Cps14J of S. pneumoniae (20). Cps2J showed similarity with Cps14I of S. pneumoniae as well. Cps14I was shown to have N-acetyl glucosaminyltransferase activity, whereas Cps14J has a β-1, 4-galactosyltransferase activity (20). In S. pneumoniae, Cps14I is responsible for the addition of the third sugar and Cps14J for the addition of the last sugar in the synthesis of the type 14 repeating unit (20). Because the capsule of S. suis type 2 contains galactose as well as N-acetyl glucosamine components, galactosyltransferase as well as N-acetyl glucoaminyltransferase activities could be envisaged for the cps2J and cps2K gene products, respectively. As was observed for Cps14I and Cps14J, the N-termini of Cps2J and Cps2K showed a significant degree of sequence similarity. Within the N-terminal domains of Cps14I and Cps14J, two small regions were identified, which were also conserved in several other glycosyltransferases (22). Within these two regions, two Asp residues were proposed to be important for catalytic activity. The two conserved regions, DXS and DXDD, were also found in Cps2J and Cps2K.

[0139] The function of Cps2I remains unclear. Cps2I showed some similarity with a protein of A. actinomycetemcomitans. Although this protein part is of the gene cluster responsible for the serotype-B-specific antigens, the function of the protein is unknown.

specific for *S. suis* serotypes 1, 2 and 9. After the entire *cps2* locus of *S. suis* serotype 2 was cloned and characterized, functions for most of the *cps2* gene products could be assigned by sequence homologies. Based on these data, the glycosyltransferase activities, required for type specificity, could be located in the center of the operon. Cross-hybridization experiments, using the individual *cps2* genes as probes on chromosomal DNAs of the 35 different serotypes, confirmed this idea. The regions containing the type-specific genes of serotypes 1 and 9 could be cloned and characterized, showing that an identical genetic organization of the *CpS* operons of other *S. suis* serotypes exists. The *cps1E*, *cps1F*, *cps1G*, *cps1H*, and *cps1I* genes revealed a

striking similarity with [cps14 E]cps14E, cps14F, cps14G, cps14H and cps14J genes of S. pneumoniae. Interestingly, S. pneumoniae serotype 14 is the serotype most commonly associated with pneumococcal infections in young children (54), whereas S. suis serotype 1 strains are most commonly isolated from piglets younger than 8 weeks (46). In S. pneumoniae, the cps14E, cps14G, cps14I and cps14J encode the glycosyltransferases required for the synthesis of the type 14 tetrameric repeating unit, showing that the cps1E, cps1G and cps1I genes encoded glycosyltransferases. The precise functions of these genes as well as the substrate specificities of the enzymes can be established. In S. pneumoniae, the cps14E gene was shown to encode a glucosyl-1-phosphate transferase catalyzing the transfer of glucose to a lipid carrier. Moreover, cpsE-like genes were found in S. pneumoniae serotypes 9N, 13, 14, 15B, 15C, 18F, 18A and 19F (60). CpsE mutants were constructed in the serotypes 9N, 13, 14 and 15B. All mutant strains lacked glucosyltransferase activity (60). Moreover, in all these S. pneumoniae serotypes, the cpsE gene seemed to be responsible for the addition of glucose to the lipid carrier. Based on these data, we suggest that in S. suis type 1, the cps IE gene may fulfil a similar function. The structure of the S. suis type 1 capsule is unknown, but it is composed of glucose, galactose, N-acetyl glucosamine, N-acetyl galactosamine and sialic acid in a ratio of 1: 2.4: 1: 1:1.4 (5). Therefore, a role of a cpsElike glucosyltransferase activity can easily be envisaged. [CpsE like] CpsE-like sequences were also found in serotypes 2, 1/2 and 14.

[0141] For polysaccharide biosynthesis in *S. pneumoniae* type 14, transfer of the second sugar of the repeating unit to the first lipid-linked sugar is performed by the gene products of cps14F and cps14G (20). Similar to Cps14F and Cps14G, the *S. suis* type 1 proteins Cps1F and Cps1G may act as one glycosyltransferase performing the same reaction. Cps14F and Cps14G of *S. pneumoniae* showed similarity to the N-terminal half and C-terminal half of the SpsK protein of Sphingomonas (20, 67), respectively. This suggests a combined function for both proteins. Moreover, cps14F\_and cps14G\_like sequences were found in several serotypes of *S. pneumoniae* and these genes always seemed to exist together (60). The same was observed for *S. suis* type 1. The cps1F and cps1G probes hybridized with type 1 and type 14 strains.

[0142] According to the similarity found between the *cps1H* gene and the *cps14H* gene of *S. pneumoniae* (20), *cps1H* is expected to encode a polysaccharide polymerase.

[0143] The protein encoded by the *cps1I* gene showed some similarity with the Cps14J protein of *S. pneumoniae* (19). The *cps14J* gene was shown to encode a β-1, 4-galactosyltransferase activity, responsible for the addition of the fourth (*i.e.* last) sugar in the synthesis of the *S. pneumoniae* serotype 14 polysaccharide. In *S. suis* type 2, the proteins encoded by the *cps2J* and *cps2K* genes showed similarity to the Cps14J protein. However, no significant homologies were found between Cps2J, Cps2K and Cps1I. In the N-terminal regions of Cps14J and Cps14I, two small conserved regions, DXS and DXDD, were identified (19). These regions seemed to be important for catalytic activity (13). At the same positions in the sequence, Cps2I contained the regions DXS and DXED.

[0144] In the region between Cps1G and Cps1H, three small Orfs were identified. Since the Orfs were expressed in three different reading frames, and did not contain potential start sites, expression is not expected. However, the three potential gene products encoded by this region showed some similarity with three successive regions of the C-terminal part of the EpsK protein of Streptococcus thermophilus (27% identity, 40). The region related to the first 82 amino acids is lacking. The EpsK protein was suggested to play a role in the export of the exopolysaccharide by rendering the polymerized exopolysaccharide more hydrophobic through a lipid modification. These data could suggest that the sequences in the region between Cps1G and Cps1H originated from epsK-like sequence. Hybridization experiments showed that this epsK-like region is also present in other serotype 1 strains as well as in serotype 14 strains (results not shown).

[0145] The function of most of the cloned serotype 9 genes can be established. Based on sequence similarity data, the *cps9E* and *cps9F* genes could be glycosyltransferases (61, 24, 63, 64, 65). Moreover, the *cps9G* and *cps9H* genes showed similarity to genes located in regions involved in polysaccharide biosynthesis, but the function of these genes is unknown (68).

[0146] Cross-hybridization experiments using the individual cps2, cps1 and cps9 genes as probes[,] showed that the cps9G and cps9H probes specifically hybridized with serotype 9 strains.

[0147] Therefore, these are useful as tools for the identification of *S. suis* type 9 strains both for diagnostic purposes as well as in epidemiological and transmission studies. We previously

developed a PCR method which can be used to detect *S. suis* strains in nasal and tonsil swabs of pigs (62). The method was used to identify pathogenic (EF-positive) strains of *S. suis* serotype 2. Besides *S. suis* type 2 strains, serotype 9 strains are frequently isolated from organs of diseased pigs. However, until now, a rapid and sensitive diagnostic test was not available for type 9 strains. Therefore, the type 9 specific probes or the type 9 specific PCR is of great diagnostic value. The *cps1F*, *cps1G* and *cps1I* probes hybridized with serotype 1 as well as with serotype 14 strains. In coagglutination tests, type 1 strains react with the anti-type 1 as well as with the anti-type 14 antisera (56). This suggests the presence of common epitopes between these serotypes. On the other hand, type 1 strains agglutinated only with anti-type 1 serum (56, 57), indicating that it is possible to detect differences between those serotypes.

[0148] The cps2F, cps2G, cps2H, cps2I and cps2J probes hybridized with serotypes 2 and 1/2 only. Serotype 34 showed a weak hybridizing signal with the cps2G probe. As shown in agglutination tests, type 1/2 strains react with sera directed against type 1 as well as with sera directed against type 2 strains (46). Therefore, type 1/2 shared antigens with both types 1 and 2. Based on the hybridization patterns of serotype 1/2 strains with the cps1 and cps2 specific genes, serotype 1/2 seemed to be more closely related to type 2 strains than to type 1 strains. In our current studies, we identify type-specific genes, primers or probes which are used for the discrimination of serotypes 1, 14 and 2 and 1/2 and others of the 35 serotypes yet known. Furthermore, type-specific genes, primers or probes can now easily be developed for yet unknown serotypes, once they become isolated.

#### Cloning and characterization of a further part of the cps2 locus.

[0149] Based on the established sequence, 11 genes, designated cps2L to cps2T, orf2U and orf2V, were identified. A gene homologous to genes involved in the polymerization of the repeating oligosaccharide unit (cps2O) as well as genes involved in the synthesis of sialic acid (cps2P to cps2T) were identified. Moreover, hybridization experiments showed that the genes involved in the sialic acid synthesis are present in *S. suis* serotypes 1, 2, 14, 27 and 1/2. The "cps2M" and "cps2N" regions showed similarity to proteins involved in the polysaccharide biosynthesis of other [g]Gram-positive bacteria. However, these regions seemed to be truncated

or were nonfunctional as the result of frame-shift or point mutations. At its 3'-end, the cps2 locus contained two insertional elements ("orf2U" and "orf2V"), both of which seemed to be nonfunctional.

[0150] To clone the remaining part of the cps2 locus, sequences of the 3'-end of pCPS26 (FIG. 1, part C) were used to identify a chromosomal fragment containing cps2 sequences located further downstream. This fragment was cloned in pKUN19, resulting in pCPS29. Using a similar approach, we subsequently isolated the plasmids pCPS30 and pCPS34 containing downstream cps2 sequences (FIG. 1, part C).

### Analysis of the cps2 operon.

[0151] The complete nucleotide sequence of the cloned fragments was determined. Examination of the compiled sequence revealed the presence of:[] a sequence encoding the C-terminal part of Cps2K, six apparently functional genes (designated cps2O-cps2T) and the remnants of 5 different ancestral genes (designated "cps2L", "cps2M", "cps2N", "orf2U" and "orf2V"). The latter genes seemed to be truncated or incomplete as the result of the presence of stop codons or frame-shift mutations [Fig. 1A] (FIG. 1, part A). Neither potential promoter sequences nor potential stem-loop structures could be identified within the sequenced region. A ribosome-binding site precedes each ORF and the majority of the ORFs are very closely linked. Three intergenic gaps were found: one between "cps2M" and "cps2N" (176 nucleotides), one between cps2O and cps2P (525 nucleotides), and one between cps2T and "orf2U" (200 nucleotides). These and our above data show that Orf2X and Cps2A-Orf2T are part of a single operon.

[0152] A list of all loci and their properties is shown in Table 4. The "cps2L" region contained three potential ORFs[,] of 103, 79 and 152 amino acids, respectively, which were only separated from each other by stop codons. Only the first ORF is preceded by a potential ribosomal binding site and contained a methionine start codon. This suggests that "cps2L" originates from an ancestral cps2L gene, which coded for a protein of 339 amino acids. The function of this hypothetical Cps2L protein remains unclear so far: no significant homologies were found between Cps2L and proteins present in the data libraries. It is not clear whether the first ORF of the

"cps2L" region is expressed into a protein of 103 amino acids. The "cps2M" region showed homology to the N-terminal 134 amino acids of the NeuA proteins of Streptococcus agalactiae and Escherichia coli (AB017355, 32). However, although the "cps2 M" region contained a potential ribosome binding site, a methionine start codon was absent. Compared with the S. agalactiae sequence, the ATG start codon was replaced by a lysin encoding AAG codon. Moreover, the region homologous to the first 58 amino acids of the S. agalactiae NeuA (identity 77%) was separated from the region homologous to amino acids 59-134 of NeuA by a repeated DNA sequence of 100-bp (see, herein). In addition, the region homologous to amino acids 59 to 95 of NeuA (identity 32%) and the region homologous to the amino acids 96 to 134 of NeuA (identity 50%) were present in different reading frames. Therefore, the partial and truncated NeuA homologue is probably nonfunctional in S. suis. The "cps2N" region showed homology to CpsJ of S. agalactiae (accession no. AB017355). However, sequences homologous to the first 88 amino acids of CpsJ were lacking in S. suis. Moreover, the homologous region was present in two different reading frames. The protein encoded by the cps2O gene showed homology to proteins of several streptococci involved in the transport of the oligosaccharide repeating unit (accession no. AB017355), suggesting a similar function for Cps2O. The proteins encoded by the cps2P, cps2S and cps2T genes showed homology to the NeuB, NeuD and NeuA proteins of S. agalactiae and E. coli (accession no. AB017355). Because the "cps2M" region also showed homology to NeuA of E. coli, the S. suis cps2 locus contains a functional neuA gene (cps2T) as well as a nonfunctional ("cps2M") gene. The mutual homology between these two regions showed an identity of 77% at the amino acid level over amino acids 1-58 and 49% over the amino acids 59-134. Cps2Q and Cps2R showed homology to the N-terminal and C-terminal parts of the NeuC protein of S. agalactiae and E. coli, respectively. This suggests that the function of the S. agalactiae NeuC protein in S. suis is likely fulfilled by two different proteins. In E. coli, the neu genes are known to be involved in the synthesis of sialic acid. NeuNAc is synthesized from N-acetylmannosamine and phosphoenolpyruvate by NeuNAc synthetase. Subsequently, NeuNAc is converted to CMP-NeuNAc by the enzyme CMP-NeuNAc synthetase. CMP-NeuNAc is the substrate for the synthesis of polysaccharide. In E. coli, K1 NeuB is the NeuNAc synthetase, and NeuA is the CMP-NeuNAc synthesis. NeuC has been implicated in the NeuNAc synthesis, but

its precise role is not known. The precise role of NeuD is not known. A role of the Cps2P-Cps2T proteins in the synthesis of sialic acid can easily be envisaged, since the capsule of *S. suis* serotype 2 is rich in sialic acid. In *S. agalactiae*, sialic acid has been shown to be critical to the virulence function of the type III capsule. Moreover, it has been suggested that the presence of sialic acid in the capsule of bacteria which can cause meningitis may be important for these bacteria to breach the blood-brain barrier. So far, however, the requirement of the sialic acid for virulence of *S. suis* remains unclear.

[0153] "Orf2U" and "Orf2V" showed homology to proteins located on two different insertional elements. "Orf2U" is homologous to IS1194 of *Streptococcus thermophilus*, whereas "Orf2V" showed homology to a putative transposase of *Streptococcus pneumoniae*. This putative transposase was recently found to be associated with the type 2 capsular locus of *S. pneunioniae*. Compared with the original insertional elements in *S. thermophilus* and *S. pneumoniae*, both "Orf2U" and "Orf2V" are likely to be non[-]functional due to frame shift mutations within their coding regions.

[0154] A striking observation was the presence of a sequence of 100 bp (FIG. 10) which was repeated three times within the cps2 operon. The sequence is highly conserved (between 94% and 98%) and was found in the intergenic regions between cps2G and cps2H, within "cps2M" and between cps2O and cps2P. No significant homologies were found between this 100-bp direct repeat sequence and sequences present in the data libraries, suggesting that the sequence is unique for *S. suis*.

# Distribution of the cps2 sequences among the 35 S. suis serotypes.

[0155] To examine the presence of sialic acid encoding genes in other *S. suis* serotypes, we performed cross-hybridization experiments. DNA fragments of the individual cps2 genes were amplified by PCR, radiolabeled with 32P and hybridized to chromosomal DNA of the reference strains of the 35 different *S. suis* serotypes. As a positive control, we used a probe specific for *S. suis* 16S rRNA. The 16S rRNA probe hybridized with almost equal intensities to all serotypes tested (Table 4). The "cps2L" sequence hybridized with DNA of serotypes 1, 2, 14 and 1/2. The "cps2M", cps2O, cps2P, cps2Q, cps2R, cps2S and cps2T genes hybridized with DNA of serotypes

1, 2, 14, 27 and 1/2. Because the cps2P-cps2T genes are most likely involved in the synthesis of sialic acid, these results suggest that sialic acid is also a part of the capsule in the *S. suis* serotypes 1, 2, 14, 27 and 1/2. This is in agreement with the finding that the serotypes 1, 2 and 1/2 possess a capsule that is rich in sialic acid. Although the chemical compositions of the capsules of serotypes 14 and 27 are unknown, recent agglutination studies using sialic acid-binding lectins suggested the presence of sialic acid in *S. suis* serotype 14, but not in serotype 27. In these studies, sialic acid was also detected in serotypes 15 and 16. Since the latter observation is not in agreement with our hybridization studies, it might be that other genes, not homologous to the cps2P-cps2T genes, are responsible for the sialic acid synthesis in serotypes 15 and 16.

A probe based on "cps2N" sequences hybridized with DNA from serotypes 1, 2, [0156] 14 and 1/2. A probe specific for "orf2U" hybridized with serotypes 1, 2, 7, 14, 24, 27, 32, 34, and 1/2, whereas a probe specific for 'orf2V" hybridized with many different serotypes. In addition, we prepared a probe specific for the 100-bp direct repeat sequence. This probe hybridized with the serotypes 1, 2, 13, 14, 22, 24, 27, 29, 32, 34 and 1/2 (Table 4). To analyze the number of copies of the direct repeat sequence within the S. suis serotype 2 chromosome, a Southern blot hybridization and analysis was performed. Therefore, chromosomal DNA of S. suis serotype 2 was digested with NcoI and hybridized with a 32P-labeled direct repeat sequence. Only one hybridizing fragment, containing the three direct repeats present on the cps2 locus, was found (results not shown). This indicates that the 100-bp direct repeat sequence is only associated with the cps2 locus. In S. pneumoniae, a 115-bp long repeated sequence was found to be associated with the capsular genes of serotypes 1, 3, 14 and 19F. In S. pneumoniae, this 115-bp sequence was also found in the vicinity of other genes involved in pneumococcal virulence (hyaluronidase and neuraminidase genes). A regulatory role of the 115-bp sequence in co[-]ordinate control of these virulence- related genes was suggested.

[0157] To study the role of the capsule in resistance to phagocytosis and in virulence, we constructed two isogenic mutants in which capsule synthesis was disturbed. In 10cpsB, the cps2B gene was disturbed by the insertion of an antibiotic-resistance gene, whereas in 10cpsEF, parts of the cps2E and cps2F genes were replaced. Both mutant strains seemed to be completely unencapsulated. Because the [cps 2]cps2 genes seemed to be part of an operon, polar effects

cannot be excluded. Therefore, these data did not give any information about the role of Cps2B. Cps2E or Cps2F in the polysaccharide biosynthesis. However, the results clearly show that the capsular polysaccharide of S. suis type 2 is a surface component with antiphagocytic activity. In vitro wild type encapsulated bacteria are ingested by phagocytes at a very low frequency, whereas the mutant unencapsulated bacteria are efficiently ingested by porcine macrophages. Within 2 hours, over 99.6% of mutant bacteria were ingested and over 92% of the ingested bacteria were killed. Intracellularly, wild type as well as mutant strains seemed to be killed with the same efficiency. This suggests that the loss of capsular material is associated with loss of capacity to resist uptake by macrophages. This loss of resistance to in vitro phagocytosis was associated with a substantial attenuation of the virulence in germfree pigs. All pigs inoculated with the mutant strains survived the experiment and did not show any specific clinical signs of disease. Only some aspecific clinical signs of disease could be observed. Moreover, mutant bacteria could be reisolated from the pigs. This supports the idea that, as in other pathogenic Streptococci, the capsule of S. suis acts as an important virulence factor. Transposon mutants prepared by Charland impaired in the capsule production showed a reduced virulence in pigs and mice. To construct these mutants, the type 2 reference strain S735 was used. We previously showed that this strain is only weakly virulent for young pigs. Moreover, the insertion site of the transposon is unsolved so far.

# As a further example herein, a rapid PCT test for Streptococcus suis type 7 is described.

[0158] Recent epidemiological studies on *Streptococcus suis* infections in pigs indicated that, besides serotypes 1, 2 and 9, serotype 7 is also frequently associated with diseased animals. For the latter serotype, however, no rapid and sensitive diagnostic methods are available. This hampers prevention and control programs. Here we describe the development of a type-specific PCR test for the rapid and sensitive detection of *S. suis* serotype 7. The test is based on DNA sequences of capsular (cps) genes specific for serotype 7. These sequences could be identified by cross-hybridization of several individual cps genes with the chromosomal DNAs of 35 different *S. suis* serotypes.

[0159] Streptococcus suis is an important cause of meningitis, septicemia, arthritis and sudden death in young pigs (69, 70). It can, however, also cause meningitis in man (71). Attempts to control the disease are still hampered by the lack of sufficient knowledge about the epidemiology of the disease and the lack of effective vaccines and sensitive diagnostics.

[0160] S. suis strains can be identified and classified by their morphological, biochemical and serological characteristics (70, 73, 74). Serological classification is based on the presence of specific antigenic determinants. Isolated and biochemically characterized S. suis cells are agglutinated with a panel of specific sera. These typing methods are very laborious and time-consuming and can only be performed on isolated colonies. Moreover, it has been reported that nonspecific cross-reactions may occur among different types of S. suis (75, 76).

[0161] So far, 35 different serotypes have been described (7, 78, 79). S. suis serotype 2 is the most prevalent type isolated from diseased pigs, followed by serotypes 9[,] and 1. However, recently, serotype 7 strains were also frequently isolated from diseased pigs (80, 81, 82). This suggests that infections with S. suis serotype 7 strains seem to be an increasing problem. Moreover, the virulence of S. suis serotype 7 strains was confirmed by experimental infection of young pigs (83).

[0162] Recently, rapid and sensitive PCR assays specific for serotypes 2 (and 1/2), 1 (and 14) and 9 were developed (84). These assays were based on the cps loci of *S. suis* serotypes 2, 1 and 9 (84, 85). However, until now, no rapid and sensitive diagnostic test [is]was available for *S. suis* serotype 7. Herein we describe the development of a PCR test for the rapid and sensitive detection of *S. suis* serotype 7 strains. The test is based on DNA sequences which form a part of the cps locus of *S. suis* serotype 7. Compared with the serological serotyping methods, the PCR assay was a rapid, reliable and sensitive assay. Therefore, this test, in combination with the PCR tests which we previously developed for serotypes 1, 2 and 9, will undoubtedly contribute to a more rapid and reliable diagnosis of *S. suis* and may facilitate control and eradication programs.

#### Materials and Methods

# Bacterial strains, growth conditions and serotyping.

[0163] The bacterial strains and plasmids used in this study are listed in Table 7. The S. suis reference strains were obtained from M. Gottschalk, Canada. S. suis strains were grown in Todd-Hewitt broth (code CM189, Oxoid), and plated on Columbia agar blood base (code CM331, Oxoid) containing 6% (v/v) horse blood. E. coli strains were grown in Luria broth (86) and plated on Luria broth containing 1.5% (w/v) agar. If required, ampicillin was added to the plates. The S. suis strains were serotyped by the slide agglutination test with serotype-specific antibodies (70).

# DNA techniques.

[0164] Routine DNA manipulations and PCR reactions were performed as described by Sambrook et al. (88). Blotting and hybridization [was]were performed as described previously (84, 86).

# DNA sequence analysis.

[0165] DNA sequences were determined on a 373A DNA Sequencing System (Applied Biosystems, Warrington, GB). Samples were prepared by use of an ABI/PRISM dye terminator cycle sequencing ready reaction kit (Applied Biosystems). Custom-made sequencing primers were purchased from Life Technologies. Sequencing data were assembled and analyzed using the McMollyTetra program. The BLAST program was used to search for protein sequences homologous to the deduced amino acid sequences.

#### PCR.

[0166] The primers used for the cps7H PCR correspond to the positions 3334-3354 and 3585-3565 in the *S. suis* cps7 locus.

The sequences were:

- 5' -AGCTCTAACACGAAATAAGGC-3' (SEQ. ID. No. 7) and
- 5'-GTCAAACACCCTGGATAGCCG3' (SEQ. ID. No. 8).

The reaction mixtures contained 10 mM Tris-HC1, pH 8.3; 1.5 mM

MgC12; 50 mM KC1; 0.2 mM of each of the four deoxynucleotide triphosphates; 1 microM of each of the primers and 1U of AmpliTaq Gold DNA polymerase (Perkin Elmer Applied Biosystems, New Jersey). DNA amplification was carried out in a Perkin Elmer 9600 thermal cycler and the program consisted of an incubation for 10 min at 95°C and 30 cycles of 1 min at 95°C, 2 min at 56°C and 2 min at 72°C.

#### Results and discussion

# Cloning of the seroytpe 7-specific cps genes.

[0167] To isolate the type-specific cps genes of *S. suis* serotype 7, we used the cps9E gene of serotype 9 as a probe to identify chromosomal DNA fragments of type 7 containing homologous DNA sequences (84). A 1.6-kb PstI fragment was identified and cloned in pKUN19. This yielded pCPS7-1 (FIG. 11, part C). In turn, this fragment was used as a probe to identify an overlapping 2.7 kb ScaI-ClaI fragment. pGEM7 containing the latter fragment was designated pCPS7-2 (FIG. 11, part C).

# Analysis of the cloned cps7 genes.

[0168] The complete nucleotide sequences of the inserts of pCPS7-1, pCPS7-2 were determined. Examination of the cps7 sequence revealed the presence of two complete and two incomplete open reading frames (ORFs) (FIG. 11, part C). All ORFs are preceded by a ribosome-binding Site. In accord with the data obtained for the cps1, cps2 and cps9 genes of serotypes 1, 2 and 9, respectively, the type 7 ORFs are very closely linked to each other. The only significant intergenic gap was that found between cps7E and cps7F (443 nucleotides). No obvious promoter sequences or potential stem-loop structures were found in this region. This suggests that, as in serotypes 1, 2 and 9, the cps genes in serotype 7 form part of an operon.

[0169] An overview of the ORFs and their properties is shown in Table 8. As expected on the basis of the hybridization data (84), the Cps9E and Cps7E proteins showed a high similarity (identity 99%, Table 8). Based on sequence comparisons between Cps9E and Cps7E, the PstI fragment of pCPS7-1 lacks the region encoding the first 371 codons of Cps7E. The C-terminal part of the protein encoded by the cps7F gene showed some similarity with the Bp1G protein of

Bordetella pertussis (88), as well as with the C-terminal part of S. suis Cps2E (85). Both Bp1G and Cps2E were suggested to have glycosyltransferase activity and are probably involved in the linkage of the first sugar to the lipid carrier (85, 88). The protein encoded by the cps7G gene showed similarity with the [B1pF] Bp1F protein of Bordetella pertussis (88). Bp1F is likely to be involved in the biosynthesis of an amino sugar, suggesting a similar function for Cps7G. The protein encoded by the cps7H gene showed similarity with the WbdN protein of E. coli (89) as well as with the N-terminal part of the Cps2K protein of S. suis (81). Both WbdN and Cps2K were suggested to have glycosyltransferase activity (85, 89).

# Serotype 7 specific cps genes.

[0170] To determine whether the cloned fragments in pCPS7-1 and pCPS7-2 contained serotype 7-specific DNA sequences, cross[]-hybridization experiments were performed. DNA fragments of the individual cps7 genes were amplified by PCR, labeled with 32P, and used to probe spot blots of chromosomal DNA of the reference strains of 35 different *S. suis* serotypes. The results are summarized in Table 9. As expected, based on the data obtained with the cps9E probe (84), the cps7E probe hybridized with chromosomal DNA of many different *S. suis* serotypes. The cps7F and cps7G probes showed hybridization with chromosomal DNA of *S. suis* serotypes 4, 5, 7, 17, and 23. However, the cps7H probe hybridized with chromosomal DNA of serotype 7 only, indicating that this gene is specific for serotype 7.

# Type specific PCR.

[0171] We tested whether we could use PCR instead of hybridization for the typing of the *S. suis* serotype 7 strains. For that purpose, we selected an oligonucleotide primer set within the cps7H gene with which an amplified fragment of 251-bp was expected. In addition, we included in our analysis several *S. suis* serotype 7 strains, other than the reference strain. These strains were obtained from different countries and were isolated from different organs (Table 7). The results show that indeed a fragment of about 250-bp was amplified with all type 7 strains used (FIG. 12, part B), whereas no PCR products were obtained with serotype 1, 2 and 9 strains (FIG. 12, part A). This suggests that the PCR test, as described here, is a rapid diagnostic tool for

the identification of *S. suis* serotype 7 strains. Until now, such a diagnostic test was not available for serotype 7 Strains. Together with the recently developed PCR assays for serotypes 1, 2, 1/2, 14 and 9, this assay may be an important diagnostic tool to detect pigs carrying serotype 2, 1/2, 1, 14, 9 and 7 strains and may facilitate control and eradication programs.

| strain/plasmid  | relevant  | source/referen |
|-----------------|---|----------------|
|                 | characteristics                                   |                |
|                 |   |                |
| Strain          |   |                |
| E.coli          | a   | (28)           |
| CC118           | PhoA*   | (20)           |
| XL2 blue        | Stratagene  |                |
| E.coli          |   |                |
| XL2 blue        | Stratagene  |                |
| S. svis         |   |                |
| 10              | virulent serotype 2 strain                        | (49)           |
| 3               | serotype 2  | (63)           |
| 17              | serotype 2  | (63)           |
| 735             | reference strain serotype 2                       | (63)           |
| 115             | serotype 2  | (63)           |
| 5555            | reference strain serotype 1                       | (63)           |
| 308             | serotype 1  | (63)           |
| 290             | serotype 1  | (63)           |
| 637             | serotype 1  | (63)           |
| 673             | serotype 1/2                                      | (63)           |
| -<br>679        | serotype 1/2                                      | (63)           |
| 928             | serotype 1/2                                      | (63)           |
| 934             | serotype 1/2                                      | (63)           |
|                 | reference strains serotype 1/2                    | (63)           |
| . 18            | reference strain serotype 9                       | (63)           |
| 73              | serotype 9  | (63)           |
| 37              | serotype 9  | (63)           |
| 07              | serotype 9  | (63)           |
| ference strains | serotypes 1-34                                    | (9, 56, 14)    |
| suis            |   |                |
|                 | virulent serotype 2 strain                        | (51)           |
| psB             | isogenic cpsB mutant of strain 10                 | this work      |
| psEF            | isogenic cpsEF mutant of strain 10                | this work      |
| enid .          |   | 403)           |
| N19             | replication functions pUC, Amp <sup>R</sup>       | (23)           |
| 472£ (+)        | replication functions pUC, Amp <sup>R</sup>       | Promega Corp.  |
| L9R             | replication functions pUC, Amp <sup>A</sup>       | (29)           |
| 20R             | replication functions pUC, Amp <sup>R</sup>       | (29)           |
| spc             | pIC19R containing spc <sup>R</sup> gene of pDL282 | labcollection  |

| pDL282  | replication functions of pBR322 al                                   | (43)              |
|---------|--|-------------------|
|         | pVT736-1, Amp <sup>R</sup> , Spc <sup>R</sup>                        | this work         |
| PPHOS2  | pIC-spc containing the truncated phoA gene                           | CHIES MOLK        |
|         | of pPHO7 as a PstI-BamHI fragment                                    |                   |
| рРН07   | contains truncated phoA gene   | (15)              |
| pPHOS7  | pPHOS2 containing chromosomal S. suis DNA                            | this work         |
| pCPS6   | pKUN19 containing 6 kb HindIII fragment                              | this work (Fig.1) |
| •       | of cps operon  |                   |
| pCPS7   | pKUN19 containing 3,5 kb EcoRI-HindIII fragment                      | this work (Fig.1) |
| •       | of cps operon  |                   |
| pCPS11  | pCPS7 in which 0.4 kb PstI-BamHI fragment                            | this work (Fig.1) |
| •       | of <i>cps</i> B gene is replaced by Spc <sup>R</sup> gene of pIC-spc |                   |
| pCPS17  | pKUN19 containing 3.1 kb KpnI fragment                               | this work (Fig.1) |
| POLICE  | of cps operon  |                   |
| pCPS18  | pKUN19 containing 1.8 kb SnaBI fragment                              | this work (Fig.1) |
| polula  | of cps operon  |                   |
| pCPS20  | pKUN19 containing 3.3 kb XbaI-HindIII                                | this work (Fig.1) |
| peroze  | fragment of cps operon   |                   |
| pCPS23  | pGEM72f(+) containing 1.5 kb MluI fragment                           | this work (Fig.1) |
|         | of cps operon  |                   |
| pCPS25  | pIC20R containing 2.5 kb KpnI-Sall fragment                          | this work (Fig.1) |
| ,       | of pCPS17  |                   |
| pCPS26  | pKUN19 containing 3.0 kb HindIII fragment                            | this work (Fig.1) |
| P       | of cps operon  |                   |
| pCPS27  | pCPS25 containing 2.3 kb XbaI (blunt)-ClaI                           | this work (Fig.1) |
| •       | fragment of pCPS20   |                   |
| pCPS28  | pCPS27 containing the 1.2 kb PstI-XhoI SpcR                          | this work (Fig.1) |
| •       | gene of pIC-spc  |                   |
| pCPS29  | pKUN19 containing 2.2 kb SacI-PstI fragment                          | this work (Fig.1) |
|         | of cps operon  |                   |
| pCPS1-1 | pKUN19 containing 5 kb EcoRV fragment                                | this work (Fig.1) |
|         | of cps operon of type 1  |                   |
| pCPS1-2 | pKUN19 containing 2.2 kb HindIII fragment                            | this work (Fig.1) |
|         | of cps operon of type 1  |                   |
| pCPS9-1 | pKUN19 containing 1 kb HindIII-Xbal                                  | this work (Fig.1) |
|         | fragment of cps operon of serotype 9                                 |                   |
| pCPS9-2 | pKUN19 containing 4.0 kb XbaI-XbaI                                   | this work (Fig.1) |
| -       | fragment of cps operon of serotype 9                                 |                   |

Amp<sup>R</sup>: ampicillin resistant Spc<sup>R</sup>: spectinomycin resistant cps: capsular polysaccharide

# Table 1 continued

suis serotype 2 and similarities to gene product Properties of Orfs in the cps locus of S. other bacteria Table 2.

| ORF   | nucleotide<br>position in<br>sequence | number<br>of amino<br>acids | ້ວິວ | proposed function<br>of gene product <sup>1</sup> | similar gene product<br>(% identity)                    |
|-------|---------------------------------------|-----------------------------|------|---|---|
| Orf22 | 1 -719                                | 240                         | 44   | II broad all                                      |   |
| Orf2Y | 2079-822                              | 419                         | 38   | Transcription                                     | B. subtilis Yits (26%)                                  |
| Orf2X | 2202-2934                             | 244                         | 39   | regulation<br>Unknown                             | . subtilis ICXD (39%)                                   |
| Cps2A | 3041-4484                             | 481                         | 39   | Regulation  | H. influenzae YAAA (24%)<br>S. Dneumoniae Chaleen (200) |
| Cps2B | 4504-5191                             | 229                         | 40   | Chain length<br>determination                     |   |
| Cps2C | 5203-5878                             | 225                         | 40   | Chain length<br>determination/<br>Export          | S. pneumoniae Cps23fD (63%)                             |
| Cps2D | 5919-6648                             | 243                         | 38   | Unknown   | S. pneumoniae CosB (62%)                                |
| Cps2E | 6675-8052                             | 459                         | 33   | Glycosyltransferase                               | S. pneumoniae Cps14E (56%)                              |
| Cps2F | 8089-9256                             | 389                         | 32   | Glycosyltransferase                               | S. pneumoniae Cps23fT                                   |
| Cps2G | 9262-10417                            | 385                         | 36   | Glycosyltransferase                               | S. thermophilus EpsF (25%)                              |
| Cps2H | 10808-12176                           | 457                         | 31   | Glycosyltransferase                               | S. mutans RGPEC, M (29%)                                |
| Cps2I | 12213- 13443                          | 410                         | 29   | CP polymerase                                     | S. pneumoniae Cps23fI (48%)                             |
| Cps2J | 13583-14579                           | 332                         | 29   | Glycosyltransferase                               | S. pneumoniae Cps14J (31%)                              |
| Cps2K | 14574-15576                           | 334                         | 37   | Glycosyltransferase                               | S. pneumoniae Cps14J (40%)                              |

Table 2 continued

|             | S. agalactiae Cpsf <sup>N</sup> (77%) | E. coli NeuA , W (47%) | S. agalactiae CpsJ (43%)<br>S. agalactiae CpsK (41%) | S. agalactiae NeuB (80%)<br>E. coli NeuB (59%) | S. agalactiae NeuC <sup>*</sup> (61%)<br>E. coll NeuC <sup>*</sup> (54%) | S. agalactiae Neu $C^c$ (55%)<br>E. coli Neu $C^c$ (40%) | E. coli NeuD (32%)    | S. agalactiae CpsF (49%)<br>E.coli NeuA (34%) | S. thermophilus IS1194 (51%) | S. pneumoniae orfl (85%) |
|-------------|---------------------------------------|------------------------|--|--|--|--|-----------------------|---|------------------------------|--------------------------|
| Unknown     |                                       |                        | Repeat unit<br>transporter                           | Sialic acid synthesis                          | Sialic acid synthesis  | Sialic acid synthesis                                    | Sialic acid synthesis | CMP-NeuNAc<br>synthetase                      | Transposase                  | Transposase              |
| 37          | 38                                    | 9<br>8                 | 40   | 39   | 42   | 40   | 42                    | 40  | 42                           | 37                       |
| 103         | 1                                     | ı                      | 476  | 338  | 170  | 184  | 208                   | 395   | 168                          | 116                      |
| 15618-16635 | 16811-17322                           | 17559-18342            | 18401-19802  | 20327-21341                                    | 21355-21865  | 21933-22483  | 22501-23125           | 23136-24366                                   | 24566-25488                  | 25691-26281              |
|             | "Cps2M"                               | "Cps2N"                | . Cps20 .  | Cps2P  | Cps2Q  | Cps2R  | Cps2S                 | Cps2T   | "Orf2U"                      | "Orf2V"                  |

Predicted by sequence similarity Main Similarity refers to the amino-terminal part of the gene product

Similarity refers to the carboxy-terminal part of the gene product

ORFs between " " are truncated or non-functional as the result of frame-shift or point mutations

Table 3. Properties of Orfs in the cps genes of S. suis serotypes 1 and 9 and similarities to gene products of other bacteria

|        |                                       |               |                             |                                 |           | ٠   |   |                         |
|--------|---------------------------------------|---------------|-----------------------------|---------------------------------|-----------|---|---|-------------------------|
| ORF    | nucleotide<br>position in<br>sequence | ຶ່ນ<br>+<br>ບ | number<br>of amino<br>acids | predicted<br>mol. mass<br>(kDa) | predicted | proposed function<br>of gene product <sup>1</sup> | gene product<br>(ty)  | referenc<br>accession n |
|        |                                       |               |                             |                                 |           |   |   |                         |
| Cps1E2 | 1-1363                                | 348           | 454                         | 52.2                            | 8.0       | Glucosyltransferase                               | Streptococcus suis Cps2E  | 90                      |
| (488)  |                                       |               |                             |                                 |           |   | tococcus pneumoniae C   | (2.5)<br>ps14E<br>(12)  |
| CpslF  | 1374-1821                             | e<br>en<br>en | 149                         | 17.3                            | 8.5       | Unknown   | Streptococcus pneumoniae Cps14F<br>(83%)                                  | Cps14F<br>(14)          |
| Cps1G  | 1823-2315                             | 25.5          | 164                         | ٠.<br>د.                        | 5°.       | Glycosyltransferase                               | Streptococcus pneumoniae Cps14G(50%)                                      | Cps14G(50%)             |
| Срѕ1н  | 3035-4202                             | 24%           | . 386                       | 45.5                            | 8.<br>4.  | CP polymerase                                     | Streptococcus pneumoniae Cps14H<br>(30%)                                  | Cps14H<br>(14)          |
| Cps1I  | 4197-                                 | •             |                             |                                 |           | Glycosyltransferase                               | <u>ن</u><br>د   | Cps14J<br>(13)          |
|        |                                       |               |                             |                                 |           |   | Lactoccocus lactis EpsG (29) (31%) Streptoccoccus thermophilus EpsI (33%) | (29)<br>is Epsī<br>(28) |
| CpslJ  |                                       |               |                             | :                               |           | Glycosyltransferase                               | Streptococcus pneumoniae Cps14J   | Cps14J ( )              |

Table 3 continued

| (13) | Streptococcus pneumoniae Cps14J (44%) | Streptococcus suis Cps2D (89%) | Staphylococcus aureus Cap1D<br>(27%) | Staphylococcus aureus Cap5M<br>(52%) | Actinobacillus actinomycetemcomitans (43%) (ABOO2668_4) Haemophilus influenzae Lsg (43%) | Yersinia enterolitica RfbB (28%) |
|------|---------------------------------------|--------------------------------|--------------------------------------|--------------------------------------|--|----------------------------------|
|      | Glycosyltransferase                   | Unknown                        | Glycosyltransferase                  | Glycosyltransferase                  | Unknown  | Unknown                          |
|      | 7.8                                   | . 8.1                          |                                      | 8.2                                  | 0.8  | 7.2                              |
| •    | 32.5                                  | 24.9                           |                                      | 22.3                                 | 31.5   | 16.5                             |
|      | 278                                   | 215                            |                                      | 200                                  | 269  | 143                              |
|      | 37.8                                  | 378                            | •                                    | 36%                                  | æ<br>S   | 30\$                             |
|      |                                       | 1-646                          | -089                                 |                                      |  |                                  |
| •    | CPSIK                                 | Cps9D²                         | ತ<br>6 ಕರೆ<br>2 ರ                    | Cps9F                                | Cps9G  | Cps9H <sup>3</sup>               |

<sup>1</sup>Predicted by sequence similarity

N-terminal part of protein is lacking C-terminal part of protein is lacking

Table 4.

Hybridization of serotype 2 cps genes and neighboring sequences with chromosomal DNA of other serotypes

33 34 1/2

30. 31.32

18 19

10 +1

**Ф** 

serotypes

| • | orf2Z + + | +          | +        | + +      | +        | +              | +              | +                  |    |   |     | + | +  | + • | + |    |     | . + + | 0 | +   | +   | +   | + + S2 | 2T + + | + + | + + | + + + + + + + + + + + + + + + + + + + | 16SrRNA + + |
|---|-----------|------------|----------|----------|----------|----------------|----------------|--------------------|----|---|-----|---|----|-----|---|----|-----|-------|---|-----|-----|-----|--------|--------|-----|-----|---------------------------------------|-------------|
|   | +         | +          | +        | +        | +        | . +            | •              |                    | •  |   | •   |   | •  |     |   |    |     |       |   |     | : · |     |        |        |     | +1  |                                       | +           |
|   | +         | +          | +        | +        | 4        | ٠ ٠            | + +<br>د ،     |                    | •  | • | •   | • | •  |     | • | •  | •   |       | • |     |     | •   |        |        |     | 44  |                                       | +           |
|   | +         | +          | +        | +        | 4        | ٠ ﴿            | ٠ -            | •                  | •  | • | ٠   | ٠ | •  |     | • | •  | •   | •     | • |     | •   | ٠   | •      | •      | •   | •   |                                       | +           |
|   | +         | +          | +        | +        | • •      | + +            | ٠ ٠            | ٠                  | •  | • | •   | • | ٠, | 1   | • | •  | •   | •     | • | • ( | •   | •   | •      | •      | +   | #   | •                                     | +           |
|   | +         | +          | +        | +        |          | ٠ ٠            | + -            | +                  | •  | • | •   | • | ,  | ,   | • | .• | •   | •     | • | •   |     | •   | ٠      | •      | •   | •   | •                                     | +           |
|   | +         | +          | +        | 4        |          | •              | +              | +                  | •  | • | •   | • |    |     |   | •  | •   | •     | • | •   |     | •   | •      | •      | •   | •   | .•                                    | <b>+</b> .  |
|   | +         | +          | +        | 4        |          | + -            | +              | +                  |    |   | •   |   |    |     |   |    |     |       |   |     |     | •   | •      | •      | •   |     | •                                     | +           |
|   | •         | . +        | +        |          | •        |                | +              | +                  |    | • | •   |   |    |     |   |    |     | •     |   |     | • • |     | •      |        |     |     |                                       | +           |
|   |           | + <i>+</i> |          |          |          |                |                |                    |    |   | •   | • | ,  |     |   |    |     |       |   |     |     |     |        |        |     |     |                                       | +           |
|   | •         | +          | • •      | •        | +        | <b>+</b><br>بد | <b>+</b><br>بو | <del>+</del><br>بد | +  | • | •   | • | •  |     |   |    | +   |       |   | •   |     | •   |        |        | •   | •   | +                                     | +           |
|   | •         | <b>+</b> + | ٠ -      | •        | +        | •              | •              | •                  | •  | • | •   |   | •  |     |   |    |     |       |   |     | ٠,  | ا ا |        |        |     |     | •                                     | +           |
|   | •         | • •        | •        | ٠        | +        | ٠              | ##             | +1                 | •  | ٠ | . • |   | •  |     |   |    |     |       |   | •   | •   | • ' |        | •      | •   |     | •                                     | +           |
|   |           | + •        | •        | +.       | +        | #              | •              | +                  | •  | • | •   |   | •  |     |   |    | . • |       |   | •   | •   | •   | •      |        | •   | +   |                                       | +           |
|   |           | + -        | +        | +        | +        | #1             | •              | +                  | ٠, | • |     | 1 | •  | •   | • | •  | •   | •     | • | •   | •   | •   | •      |        | •   | +   | •                                     | +           |
|   |           | +          | +        | +        | +        | +1             |                | +                  |    |   |     |   | •  | •   |   | •  |     |       |   |     |     |     | • ,    |        | •   | -   | •                                     | +           |
|   |           |            | +1       |          |          |                | •              |                    |    |   |     |   |    |     |   |    |     |       |   |     |     |     |        |        | •   | •   | •                                     | .+          |
|   |           |            |          |          |          |                |                |                    |    |   |     | - |    |     |   |    |     |       |   |     |     |     |        | • •    |     | •   |                                       | +           |
|   |           | •          | Ŧ        | •        | •        |                | ,              | •                  |    |   |     |   |    |     |   |    |     |       |   | . • |     |     |        |        |     | . 4 | + 1                                   | +           |
|   |           | +          | <b>*</b> | <b>*</b> | <b>*</b> | •              |                |                    |    |   |     |   |    |     |   |    |     |       |   |     |     |     |        |        |     | +   |                                       | +           |
| • |           | +          | •        | +        | <b>+</b> | •              | • •            | · 4                |    | • |     |   |    |     |   |    |     |       |   |     |     |     |        |        | . + | . , | +                                     | +           |
|   |           | •          | +        |          |          |                |                |                    |    | • |     |   |    |     |   |    |     |       |   | •   |     |     |        |        | . ' | , 7 |                                       | +           |
| • |           |            |          |          |          |                |                |                    |    |   |     |   | •  |     |   |    |     |       |   | +   |     |     |        |        | . • |     |                                       | •           |
|   |           | +          |          |          |          |                |                |                    |    |   |     |   |    |     |   |    | •   |       |   |     | •   | •   |        | •      | • ' |     |                                       |             |
|   |           | +          | .\$      | •        |          | ٠ ٠            | ٠              | •                  | ٠  | • | •   | • | •  |     |   |    |     |       |   | •   | •   | •   | •      | •      |     |     | •                                     | +           |
|   |           | +          | +        | •        | • ·      | + •            | #1             | •                  | н  | • | •   | • | •  |     |   |    |     |       |   | •   |     |     |        | •      |     | •   | •                                     | •           |
|   |           | 4          | • •      |          | ٠        | +              | +              | +                  | +  | • | •   | • | •  | •   | • | •  |     | •     | • | . • | •   | •   |        |        |     | • • | •                                     | +           |
|   |           |            | 1        | •        |          | •              | •              | •                  | •  |   | •   | • | ,  |     | • | •  |     | •     | • | •   | •   | •   | •      | •      | . 1 | + + | • •                                   | +           |
|   |           |            | •        | •        | •        | •              | #              | #1                 | •  | • | ٠   | • |    |     |   |    |     |       |   | •   |     |     | •      | •      | •   | • • | •                                     | •           |
|   |           |            |          |          |          |                |                |                    |    |   |     |   |    |     |   |    |     |       |   |     |     |     |        |        |     |     | _                                     | _           |

Table 5. Hybridization of serotypes 1 and 9  $\sigma ps$  genes with chromosomal DNA of other s. suis serotypes

| Serotype cps1E cps1F cps1G cps1H cps1I cps9E cps9F cps |          | j     |       |       |            | DNA probes |       |    | . •          |        |         |
|--|----------|-------|-------|-------|------------|------------|-------|----|--------------|--------|---------|
|  | Serotype | cpslE | cpslF | cps1G | срзін      | cps11      | cps9E | L  |              | н6 ғар | 16 FRNA |
|  | 1        | +     | +     | +     | +          | +          |       |    |              |        | 1       |
| 3<br>6<br>7<br>8<br>9<br>10<br>11<br>12<br>13<br>14<br>14<br>15<br>16<br>17<br>18<br>18<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19  | 2        | +     |       | ,     | ı          | -          | ,     |    |              | 1      | +       |
| 10<br>11<br>12<br>13<br>14<br>15<br>16<br>17<br>18<br>18<br>19<br>19<br>19<br>10<br>11<br>11<br>11<br>12<br>13<br>14<br>15<br>16<br>17<br>18<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19   | ,        |       |       |       | ,          | ı          | •     | ٠  | 1            | •      | +       |
| 10<br>11<br>12<br>13<br>14<br>15<br>16<br>17<br>18<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19   | , ,      |       |       | •     | +          | •          | +     | ٠  | 1            | •      | +       |
| 5 5 6 6 7 7 7 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9  | <b>.</b> | ı     | 1     | ı     | +          | •          | +     | •  | ٠            | •      |         |
| 6  | S        | ı     | ı     | ,     | +          | •          | •     | •  |              | •      | •       |
| 11   | 9        | •     |       | ,     | •          | •          | -     | ı  | I            | •      | +       |
| 8 9 10 11 12 13 14 14 15 15 16 17 18 18 19 19 19 19 19 19 19 19 19 19 19 19 19   | 7        | ,     |       | ,     | 4          |            |       | •  | •            | •      | +       |
| 10<br>11<br>12<br>13<br>14<br>14<br>15<br>16<br>17<br>18<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19   | œ        | 1     | ,     |       | •          | ı          | +     | •  | 1            | •      | +       |
| 11   |          |       | ı     | ,     | •          | ı          | •     | •  | •            | •      | +       |
| 10<br>11<br>12<br>13<br>14<br>14<br>15<br>16<br>17<br>18<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19   | D.       | •     |       | ı     | +          | ,          | +     | +  | +            | +      | +       |
| 11<br>12<br>13<br>14<br>14<br>15<br>16<br>17<br>18<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19<br>19   | 10       |       | ı     | ı     | +          |            | +     | +  | 1            | •      | +       |
| 12   | 11       | •     | •     |       | +          |            | +     | #1 | •            | •      | +       |
| 13   | 12       |       | •     | ı     | #1         | •          | +     | #  | •            | ı      | +       |
| 14       +       +       +       +       +       +       +       +       +       11       11       12       12       13       14       14       14       15       16       17  | 13       | •     | •     | ı     | . <b>+</b> | 1          | +     | ı  | ı            | 1      | +       |
| 15       -   | 14       | +     | +     | +     | +          | +          |       | •  | 1            | ı      | +       |
| 16 17 18 19 19 20 20 21 21 21 22 22 23 24 25 26 27 28 28 28 29 29 20 20 20 20 20 20 20 20 20 20 20 20 20   | 15       | •     | 1     | •     | •          | •          | •     | •  |              | 1      | +       |
| 17       -       -       +       -   | 16       | •     | ı     |       | •          | •          | •     | •  |              | 1      | +       |
| 18   | 17       | ,     | •     | •     | +          |            | +     | •  |              | 1      | +       |
| 19   | 18       | •     | •     | •     | +          | •          | +     | 1  | . <b>•</b> , | •      | +       |
| 20   | 19       | •     | •     | •     | +          | •          | +     | •  | •            | •      | +       |
| 21 + + + + + + + + + + + + + + + + +   | 20       | ı     | ı     |       | •          | •          | •     | •  |              | •      | +       |
| 22   | 21       | •     | •     | i     | +          | •          | +     | +1 | ١.           | •      | +       |
|  | 22       | •     | 1     | ι     | •          | ,          | •     | •  | •            | 1      | +       |

Table 5 continued

TABLE 6. Virulence of wild type and capsular mutant 8. suis strains in germfree pigs

| S. suis  | 7 00 10      |                |                |                               |                                    |                             |                          |        |         |  |
|----------|--------------|----------------|----------------|-------------------------------|------------------------------------|-----------------------------|--------------------------|--------|---------|--|
| strains! | group<br>(n) | mortality* [%] | morbidity³ [8] | clinical ind<br>group         | clinical index of the group        | fever<br>index <sup>7</sup> | leuco-<br>cyte<br>index* | isolai | tion of | isolation of S. suis in pigs<br>[n] per group in |
|          |              |                |                | spec<br>symptoms <sup>5</sup> | non-spec.<br>symptoms <sup>6</sup> |                             |                          | CNS    | Serosae | joints   |
| 10       | 4            | 100            | 100            | 11                            | 88                                 | 43                          | 44                       | 2      | -       |  |
| 10cpsB   | 4            |                | . 0            | o                             | 10                                 | -                           | m                        |        | o m     |  |
| 10cpsEF  | 4            | o              | 0              | • • •                         | . 0                                | #4                          |                          | -      | ı m     |  |
|          |              |                |                |                               |                                    |                             |                          |        |         |  |

<sup>1</sup> strain10 in the wild type strain, strains 10cpsB and 10cpsEF are isogenic capsular mutant strains

<sup>2</sup> piglets which died spontaneously or had to be killed for animal welfare reasons

only considering pigs with specific symptoms

clinical index: % of observations which matched the described criteria

<sup>5</sup> specific symptoms: ataxia, lameness on at least one joint, stiffness

6 non-specific symptoms: inappetance, depression

 $^{7}$  s of observations in the experimental group with a body temperature > 40 $^{\circ}$  C

 $^{9}$  % of blood samples in the group in which number of granulocytes >  $10^{10}/1$ 

Table 7.

Bacterial strains and plasmids

| relevant characteristics |                                     | serotypes 1-34 serotype 7, tonsil (1993) serotype 7, organs (1994) serotype 7, brains (1994) serotype 7 (1994) serotype 7 (1994) serotype 7, lungs (1996) serotype 7, joints (1996) serotype 7, joints (1996) serotype 7, meninges (1998) | Plasmid  pKUN19replication functions pUC, Amp <sup>R</sup> pGEM72f(+)  pCPS9-1  fragment of cps operon of serotype 9  pCPS9-2  pKUN19 containing 4.0 kb xbaI-xbaI  fragment of cps operon of serotype 9  pCPS7-1  pCPS7-1  pCPS7-2  pGEM7 containing 1.6-kb PstI fragment  of cps operon of type 7  pGEM7 containing 2.7-kb ScaI-ClaI fragment  of cps operon of type 7 |
|--------------------------|-------------------------------------|---|---|
| strain/plasmid           | <b>Strain</b><br>E.coli<br>XL2 blue | S. suis<br>reference strains<br>5667<br>7037<br>7044<br>7068<br>7646<br>7744<br>7759<br>8169<br>15913   | Plasmid pKUN19replication pGEM7zf(+) pCPS9-1 pCPS9-2 pCPS7-1  |

'Amp<sup>R</sup>; ampicillin resistant cps: capsular polysaccharide

Table 8. Properties of Orfs in the

| simil | arities to gen                        | similarities to gene products of other bacteria | similarities to gene products of other bacteria                            |
|-------|---------------------------------------|---|--|
| 0r f  | nucleotide<br>position in<br>sequence | proposed function of gene product               | similar gene product<br>(% identity)                                       |
|       |                                       |   |  |
| Cps7E | 1-719                                 | Glycosyltransferase                             | Streptococcus suis Cps9E (99%)   |
| Cps7F | 1164-1863                             | Glycosyltransferase                             | Bordetella pertussis $BplG^1$ (43%) Streptococcus suis $Cps2E^1$ (33%)     |
| Cps7G | 1872-3086                             | Biosynthesis amino sugar                        | Bordetella pertussis BplF (48%)  |
| Срз7н | 3104-3737                             | Glycosyltransferase                             | Escherichia coli WbdN (35%)<br>Streptococcus suis Cps2K <sup>2</sup> (31%) |
| •     |                                       |   | -  |

'similarity refers to the C-terminal part of the gene product 'similarity refers to the N-terminal part of the gene product

| Table 9.                                    | İ | į | ##  | Hybridization of serotype 7 cps probes with chromosomal DNA of S. suis serotypes | ridiza | atti | e    | E E | erol | ton of serotype 7 cps probes with chromosomal DNA of S. suis serotypes          | 7 6       | SQ  | prot         | 8 9          | 뒫 | d d      | TOBO           | osom - | 18        | <b>S</b> | a d       | S.        | uis | 30I | oty | <b>6</b> |           |           |   |   |         |
|---|---|---|-----|--|--------|------|------|-----|------|---|-----------|-----|--------------|--------------|---|----------|----------------|--------|-----------|----------|-----------|-----------|-----|-----|-----|----------|-----------|-----------|---|---|---------|
| serotypes                                   | - | ~ | 6   | 4  | ın     |      | _    | 8   | _    | 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 10 | 12        | 13  | <del>4</del> | <del>1</del> | ₽ | <b>1</b> | . <del>L</del> | 2      | 7         | - 2      | 23        | 24        | 23  | 8   | 27  | 28       | . 62      | 0 31      | 3 | 8 | 1 48    |
| DNA probes                                  |   |   |     |  |        |      |      |     |      |   |           |     |              |              |   |          | 1              | 1      |           |          |           |           |     |     |     |          |           |           |   |   |         |
| cps7E<br>cps7F<br>cps7G<br>cps7H<br>16S/RNA | + | + | + + | +++++  | +++,+  | +    | ++++ |     | * *  | + + + + +   | + + + + + | + + | +            | +            | + | +++.+    | + + + +        | + +    | + 1 1 1 + | +        | + + + , + | + + + + + | +   | •   | 4   |          | + 1 1 1 1 | + + + + + |   |   | 1 1 1 1 |

Table 9.

# **SEQUENCE LISTING**

- <110> Smith, Hilda
- <120> STREPTOCOCCUS SUIS VACCINES AND DIAGNOSTIC TESTS
- <130> 2183-4726
- <150> PCT/NL99/00460
- <151> 1999-07-19
- <150> EP98202465.5
- <151> 1998-07-22
- <150> EP98202467.1
- <151> 1998-07-22
- <160> 53
- <170> PatentIn version 3.0
- <210> 1
- <211> 23

| <212>  | DNA                 |    |
|--------|---------------------|----|
| <213>  | Artificial          |    |
| <220>  |                     |    |
| <223>  | Primer              |    |
| <400>  | 1                   |    |
| caaacg | caag gaattacggt atc | 23 |
|        |                     |    |
| <210>  | 2                   |    |
| <211>  | 23                  |    |
| <212>  | DNA                 |    |
| <213>  | Artificial          |    |
| <220>  |                     |    |
| <223>  | primer              |    |

<210> 3

<400> 2

gagtatctaa agaatgccta ttg

- <211> 20
- <212> DNA
- <213> Artificial
- <220>
- <223> primer
- <400> 3
- ggcggtctag cagatgctcg

20

- <210> 4
- <211> 19
- <212> DNA
- <213> Artificial
- <220>
- <223> primer
- <400> 4
- gcgaactgtt agcaatgac

- <210> 5
- <211> 21
- <212> DNA
- <213> Artificial
- <220>
- <223> primer
- <400> 5
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- <210> 6
- <211> 20
- <212> DNA
- <213> Artificial
- <220>
- <223> primer
- <400> 6

# cggaagtatc tgggctactg

20

- <210> 7
- <211> 21
- <212> DNA
- <213> Artificial
- <220>
- <223> primer
- <400> 7
- agctctaaca cgaaataagg c
- <210> 8
- <211> 21
- <212> DNA
- <213> Artificial
- <220>
- <223> primer

<400> 8

gtcaaacacc ctggatagcc g

21

- <210> 9
- <211> 6992
- <212> DNA
- <213> Streptococcus suis
- <220>
- <221> misc\_feature
- <222> (1)..(6992)
- <223> CPS 2
- <400> 9

atcgccaaac gaaattggca ttatttgata tgatagcagt tgcaatttct gcaatcttaa 60

caagtcatat accaaatgct gatttaaatc gttctggaat ttttatcata atgatggttc 120

attattttgc attttttata tctcgtatgc cagttgaatt tgagtataga ggtaatctga 180

tagagtttga aaaaacattt aactatagta taatatttgc aatttttctt acggcagtat 240

catttttgtt ggagaataat ttcgcacttt caagacgtgg tgccgtgtat ttcacattaa 300

taaacttcgt tttggtatac ctatttaacg taattattaa gcagtttaag gatagctttc 420 tattttcgac aatctatcaa aaaaagacga ttctaattac aacggctgaa cgatgggaaa atatgcaagt tttatttgaa tcacataaac aaattcaaaa aaatcttgtt gcattggtag 480 540 ttttaggtac agaaatagat aaaattaatt tatcattacc gctctattat tctgtggaag 600 aagctataga gttttcaaca agggaagtgg tcgaccacgt ctttataaat ctaccaagtg 660 agtttttaga cgtaaagcaa ttcgtttcag attttgagtt gttaggtatt gatgtaagcg 720 ttgatattaa ttcattcggt tttactgcgt tgaaaaacaa aaaaatccaa ctgctaggtg accatagcat tgtaactttt tccacaaatt tttataagcc tagtcatatc atgatgaaac 780 gacttttgga tatactcgga gcggtagtcg ggttaattat ttgtggtata gtttctattt 840 tgttagttcc aattattcgt agagatggtg gaccggctat ttttgctcag aaacgagttg 900 gacagaatgg acgcatattt acattctaca agtttcgatc gatgtatgtt gatgctgagg 960 agcgcaaaaa agacttgctc agccaaaacc agatgcaagg gtgggtatgt tttaaaatgg 1020 gaaaaacgat cctagaatta ctccaattgg acatttcata cgcaaaaaca agtttagacg 1080 agttaccaca gttttataat gttttaattg gcgatatgag tctagttggt acacgtccac 1140 ctacagttga tgaatttgaa aaatatactc ctggtcaaaa gagacgattg agttttaaac 1200

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aaaaaagtat ggtgaacatg taaatgatca tcaagtagag tttgtaagaa gaattttaca 2160 agataataat attttattta tagaaaatat agatgatttg tttgaaaaaa ttattgaagt 2220 ttctaagcaa actaacttta catcaaataa taatttttt tgtgaaagat taaaacaaat 2280 agttgaaaaa tttaatgagg atcaagaaaa tgaataataa aaaagatgca tatttgataa 2340 tggcttatca taatttttct cagattttac tggagaggga tacagatatt atcatcttct 2400 ctcaggagaa tgcacaccat tagttccttc agaatacctg tataattatt ttaaatattc 2460 tcaggattta tatgttgaat ttacaaaaga tgagcaaaaa tataaagaaa ataggatata 2520 tgaacgagtt aaatgttaca gattatttcc taatatatca gaaaaaaacta ttgataatgt 2580 actgtttaga attttattaa gaatgtatcg agcttttgaa tactatttac aaagattgtt 2640 gtttattgat agaataaaaa acatggtcta agaataagat ttggttctaa ttgggtttcg 2700 cttccacatg attttgtggc aattetttta tcaaatgaaa acgaaacage ttatttattt 2760 aagtaatcta aatgtccaga tgaactattt atacagacaa ttatagaaaa atatgaattt 2820 tcaaatagat tatctaaata tggaaattta agatatataa agtggaaaaa atcaacatct 2880 tctcctattg tctttacaga tgattctatt gatgaattgc taaatgcaag aaatttaggt 2940 tttttatttg ctagaaagtt aaaaatagaa aataaatcta aatttaaaga aattattact 3000

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gtaaaaagct gctagaggcg gatggtcatc gctttgtggt ggcctgtaat aaactctata 6660

aaaaaagaact atttgaagat tttcgatttg aaaagggtaa gattcatgaa gatgaatact 6720

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gcctactgga atttcaaaat gaacgaatgg acttctatga aagtagagga gataaagagc 6900

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atcattggtt gagcaaacag caaaagaagc tt 6992

- <210> 10
- <211> 239
- <212> PRT
- <213> Streptococcus suis
- <220>
- <221> misc\_feature
- <223> ORF2Z

# <400> 10

Ser Leu Asp Ile Asp His Met Met Glu Val Met Glu Ala Ser Lys Ser Ala Ala Gly Ser Ala Cys Pro Ser Pro Gln Ala Tyr Gln Ala Ala Phe Glu Gly Ala Glu Asn Ile Ile Val Val Thr Ile Thr Gly Gly Leu Ser Gly Ser Phe Asn Ala Ala Arg Val Ala Arg Asp Met Tyr Ile Glu Glu His Pro Asn Val Asn Ile His Leu Ile Asp Ser Leu Ser Ala Ser Gly Glu Met Asp Leu Leu Val His Gln Ile Asn Arg Leu Ile Ser Ala Gly Leu Asp Phe Pro Gln Val Val Glu Ala Ile Thr His Tyr Arg Glu His Ser Lys Leu Phe Val Leu Ala Lys Val Asp Asn Leu Val Lys Asn 

Gly Arg Leu Ser Lys Leu Val Gly Thr Val Val Gly Leu Leu Asn Ile

| Arg Met Val   | Gly Glu Ala Se          | er Ala Glu G | ly Lys Leu Gl         | u Leu Leu Gln |
|---------------|-------------------------|--------------|-----------------------|---------------|
| 145           | 150                     | 155          | 160                   |               |
| Lys Ala Arg ( | Gly His Lys Ly<br>5 170 |              | nr Ala Ala Phe<br>175 | Glu Glu Met   |
| Lys Lys Ala ( | Gly Tyr Asp Gl          | y Gly Arg Il | e Val Met Ala         | His Arg Asn   |
| 180           | 185                     | 19           |                       | •             |
|               |                         |              |                       |               |
| Asn Ala Lys l | Phe Phe Gln Gl          | n Phe Ser G  | lu Leu Val Ly         | s Ala Ser Phe |
| 195           | 200                     | 205          |                       |               |
| Pro Thr Ala V | Val Ile Asp Glu         | Val Ala Thi  | r Ser Gly Leu         | Cys Ser Phe   |
| 210           | 215                     | 220          |                       |               |
|               |                         |              |                       |               |
| Tyr Ala Glu ( | Glu Gly Gly Le          | u Leu Met C  | Bly Tyr Glu Va        | ıl Lys Ala    |
| 225           | 230                     | 235          |                       |               |
|               |                         |              |                       |               |
| <210> 11      |                         |              |                       |               |
|               |                         |              |                       |               |
| <211> 244     |                         |              |                       |               |
|               |                         |              |                       |               |
| <212> PRT     |                         |              |                       |               |

<213> Streptococcus suis

<221> misc\_feature

<220>

#### <223> ORF2X

<400> 11

Met Lys Ile Ile Pro Asn Ala Lys Glu Val Asn Thr Asn Leu Glu

1 5 10 15

Asn Ala Ser Phe Tyr Leu Leu Ser Asp Arg Ser Lys Pro Val Leu Asp
20 25 30

Ala Ile Ser Gln Phe Asp Val Lys Lys Met Ala Ala Phe Tyr Lys Leu 35 40 45

Asn Glu Ala Lys Ala Glu Leu Glu Ala Asp Arg Trp Tyr Arg Ile Arg
50 55 60

Thr Gly Gln Ala Lys Thr Tyr Pro Ala Trp Gln Leu Tyr Asp Gly Leu 65 70 75 80

Met Tyr Arg Tyr Met Asp Arg Gly Ile Asp Ser Lys Glu Glu Asn 85 90 95

Tyr Leu Arg Asp His Val Arg Val Ala Thr Ala Leu Tyr Gly Leu Ile
100 105 110

His Pro Phe Glu Phe Ile Ser Pro His Arg Leu Asp Phe Gln Gly Ser 115 120 125

| Leu Lys | Ile Gly | Asn Gln | Ser Leu | Lys Gln | Tyr | Trp | Arg | Pro | Tyr | Туг |
|---------|---------|---------|---------|---------|-----|-----|-----|-----|-----|-----|
| 130     |         | 135     | 1       | 140     |     |     |     |     |     |     |

Ser Ile Lys Met

| <213> Streptococcus su | iis |
|------------------------|-----|
|------------------------|-----|

Lys Phe Ser Thr Arg Leu Asn Ser Asn Ser Thr Phe Ser Glu Tyr Glu
100 105 110

Met Ser Ile Leu Val Pro Ala Asn Ser Asp Ile Thr Asp Val Arg Gln
115 120 125

Leu Thr Ser Ile Leu Ala Pro Ala Glu Tyr Asp Gln Asp Asn Ile Thr
130 135 140

Ala Leu Leu Asp Asp Ile Ser Lys Met Glu Ser Thr Gln Leu Ala Thr
145 150 155 160

Ser Pro Gly Thr Ser Tyr Leu Thr Ala Tyr Gln Ser Met Leu Asn Gly
165 170 175

Glu Ser Gln Ala Met Val Phe Asn Gly Val Phe Thr Asn Ile Leu Glu
180 185 190

Asn Glu Asp Pro Gly Phe Ser Ser Lys Val Lys Lys Ile Tyr Ser Phe 195 200 205

Lys Val Thr Gln Thr Val Glu Thr Ala Thr Lys Gln Val Ser Gly Asp
210 215 220

Ser Phe Asn Ile Tyr Ile Ser Gly Ile Asp Ala Tyr Gly Pro Ile Ser 225 230 235 240

Thr Val Ser Arg Ser Asp Val Asn Ile Ile Met Thr Val Asn Arg Ala 245 250 255 Thr His Lys Ile Leu Leu Thr Thr Pro Arg Asp Ser Tyr Val Ala 260 265 270

Phe Ala Asp Gly Gln Asn Gln Tyr Asp Lys Leu Thr His Ala Gly 275 280 285

Ile Tyr Gly Val Asn Ala Ser Val His Thr Leu Glu Asn Phe Tyr Gly
290 295 300

Ile Asp Ile Ser Asn Tyr Val Arg Leu Asn Phe Ile Ser Phe Leu Gln
305 310 315 320

Leu Ile Asp Leu Val Gly Gly Ile Asp Val Tyr Asn Asp Gln Glu Phe 325 330 335

Thr Ser Leu His Gly Asn Tyr His Phe Pro Val Gly Gln Val His Leu
340 345 350

Asn Ser Asp Gln Ala Leu Gly Phe Val Arg Glu Arg Tyr Ser Leu Thr 355 360 365

Gly Gly Asp Asn Asp Arg Gly Lys Asn Gln Glu Lys Val Ile Ala Ala 370 375 380

Leu Ile Lys Lys Met Ser Thr Pro Glu Asn Leu Lys Asn Tyr Gln Ala 385 390 395 400

Ile Leu Ser Gly Leu Glu Gly Ser Ile Gln Thr Asp Leu Ser Leu Glu
405 410 415

Thr Ile Met Ser Leu Val Asn Thr Gln Leu Glu Ser Gly Thr Gln Phe

420

425

430

Thr Val Glu Ser Gln Ala Leu Thr Gly Thr Gly Arg Ser Asp Leu Ser
435 440 445

Ser Tyr Ala Met Pro Gly Ser Gln Leu Tyr Met Met Glu Ile Asn Gln 450 455 460

Asp Ser Leu Glu Gln Ser Lys Ala Ala Ile Gln Ser Val Leu Val Glu 465 470 475 480

Lys

<210> 13

<211> 229

<212> PRT

<213> Streptococcus suis

<220>

<221> misc\_feature

<223> CPS2B

<400> 13

| Met Asn As              | sn Gln Glu Va | l Asn Ala I | le Glu Ile Asp   | Val Leu Phe Leu   |
|-------------------------|---------------|-------------|------------------|-------------------|
| 1                       | 5             | 10          | 15               |                   |
|                         |               |             |                  |                   |
| Leu Lys Th              | r Ile Trp Arg | Lys Lys Ph  | e Leu Ile Leu I  | Leu Thr Ala Val   |
| 20                      | 25            |             | 30               |                   |
|                         |               |             |                  |                   |
| Leu Thr Ala             | a Gly Leu Ala | Phe Val Ty  | yr Ser Ser Phe   | Leu Val Thr Pro   |
| 35                      | 40            | 45          |                  |                   |
|                         |               |             |                  |                   |
| Gln Tyr As              | p Ser Thr Thr | Arg Ile Ty  | r Val Val Ser C  | In Asn Val Glu    |
| 50                      | 55            | 60          |                  |                   |
|                         |               |             |                  |                   |
| Ala Gly Ala             | Gly Leu Thr   | Asn Gln G   | lu Leu Gln Ala   | Gly Thr Tyr Leu   |
| 65                      | 70            | 75          | 80               |                   |
|                         |               |             |                  |                   |
| Ala Lys As <sub>l</sub> | p Tyr Arg Glu | Ile Ile Leu | Ser Gln Asp V    | al Leu Thr Gln    |
| 8                       | 5 9           | 90          | 95               |                   |
|                         |               |             |                  |                   |
| Val Ala Thi             | r Glu Leu Asn | Leu Lys G   | lu Ser Leu Lys   | Glu Lys Ile Ser   |
| 100                     | 10            | 5           | 110              | ·                 |
|                         |               |             |                  |                   |
| Val Ser Ile             | Pro Val Asp 7 | Thr Arg Ile | Val Ser Ile Ser  | Val Arg Asp       |
| 115                     | 120           |             | 25               |                   |
| 110                     |               |             |                  |                   |
| Ala Asn Pro             | a Asn Glu Ala | Ala Ara Ila | a Ala Asn Sar I  | Leu Arg Thr Phe   |
|                         |               | _           | c Ala Asii Sci I | Sed Aig III I IIC |
| 130                     | 135           | 140         |                  |                   |
|                         |               |             |                  |                   |

Ala Val Gln Lys Val Val Glu Val Thr Lys Val Ser Asp Val Thr Thr

Leu Glu Glu Ala Val Pro Ala Glu Glu Pro Thr Thr Pro Asn Thr Lys

165

170

175

Arg Asn Ile Leu Leu Gly Leu Leu Ala Gly Gly Ile Leu Ala Thr Gly

180

185

190

Leu Val Leu Val Met Glu Val Leu Asp Asp Arg Val Lys Arg Pro Gln

195

200

205

Asp Ile Glu Glu Val Met Gly Leu Thr Leu Leu Gly Ile Val Pro Asp

210

215

220

Ser Lys Lys Leu Lys

225

<210> 14

<211> 225

<212> PRT

<213> Streptococcus suis

<220>

<221> misc feature

<223> CPS2C

## <400> 14

Met Ala Met Leu Glu Ile Ala Arg Thr Lys Arg Glu Gly Val Asn Lys Thr Glu Glu Tyr Phe Asn Ala Ile Arg Thr Asn Ile Gln Leu Ser Gly Ala Asp Ile Lys Val Val Gly Ile Thr Ser Val Lys Ser Asn Glu Gly Lys Ser Thr Thr Ala Ala Ser Leu Ala Ile Ala Tyr Ala Arg Ser Gly Tyr Lys Thr Val Leu Val Asp Ala Asp Ile Arg Asn Ser Val Met Pro Gly Phe Phe Lys Pro Ile Thr Lys Ile Thr Gly Leu Thr Asp Tyr Leu Ala Gly Thr Thr Asp Leu Ser Gln Gly Leu Cys Asp Thr Asp Ile Pro Asn Leu Thr Val Ile Glu Ser Gly Lys Val Ser Pro Asn Pro Thr Ala 

Leu Leu Gln Ser Lys Asn Phe Glu Asn Leu Leu Ala Thr Leu Arg Arg 130 135 140

| Tyr Tyr Asp   | Tyr Val Ile Val        | Asp Cys Pro P        | ro Leu Gly Leu Val Ile  |
|---------------|------------------------|----------------------|-------------------------|
| 145           | 150                    | 155                  | 160                     |
| Asp Ala Ala 1 |                        |                      | a Met Val Ala Val Val   |
| Glu Ala Gly A | Asn Val Lys Cys<br>185 | Ser Ser Leu I<br>190 | Lys Lys Val Lys Glu Gln |
| 160           | 163                    | 190                  |                         |
| Leu Glu Gln   | Thr Gly Thr Pro        | Phe Leu Gly          | Val Ile Leu Asn Lys Tyr |
| 195           | 200                    | 205                  |                         |
| Asp Ile Ala T | hr Glu Lys Tyr         | Ser Glu Tyr G        | ly Asn Tyr Gly Lys Lys  |
| 210           | 215                    | 220                  |                         |
| Ala           |                        |                      |                         |
| 225           |                        |                      |                         |
| <210> 15      |                        |                      |                         |
| <211> 243     |                        |                      |                         |
| <212> PRT     |                        |                      |                         |

<213> Streptococcus suis

<221> misc\_feature

<220>

## <223> CPS2D

<400> 15

Met Ile Asp Ile His Ser His Ile Ile Phe Gly Val Asp Asp Gly Pro

1 5 10 15

Lys Thr Ile Glu Glu Ser Leu Ser Leu Ile Ser Glu Ala Tyr Arg Gln
20 25 30

Gly Val Arg Tyr Ile Val Ala Thr Ser His Arg Arg Lys Gly Met Phe 35 40 45

Glu Thr Pro Glu Lys Ile Ile Met Ile Asn Phe Leu Gln Leu Lys Glu
50 55 60

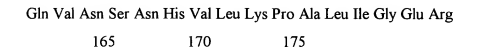
Ala Val Ala Glu Val Tyr Pro Glu Ile Arg Leu Cys Tyr Gly Ala Glu
65 70 75 80

Leu Tyr Tyr Ser Lys Asp Ile Leu Ser Lys Leu Glu Lys Lys Lys Val 85 90 95

Pro Thr Leu Asn Gly Ser Cys Tyr Ile Leu Leu Glu Phe Ser Thr Asp 100 105 110

Thr Pro Trp Lys Glu Ile Gln Glu Ala Val Asn Glu Met Thr Leu Leu 115 120 125

| Gly Leu Thr | Pro Val Leu | Ala His Ile Glu | Arg Tyr As | p Ala Leu Ala |
|-------------|-------------|-----------------|------------|---------------|
| 130         | 135         | 140             |            |               |
|             |             |                 |            |               |
| Phe Gln Ser | Glu Arg Val | Glu Lys Leu Ile | Asp Lys Gl | y Cys Tyr Thr |
| 145         | 150         | 155             | 160        |               |
|             |             |                 |            |               |



Gln Val Gln

<210> 16

<211> 459

<212> PRT

| <213> | Streptococcus | sui |
|-------|---------------|-----|
|       |               |     |

| Ser Arg Arg Gly | Ala Val Tyr Ph | ne Thr Leu Ile As | n Phe Val Leu Val |
|-----------------|----------------|-------------------|-------------------|
| 100             | 105            | 110               |                   |

| Phe Tyr Lys Pi | ro Ser His Ile | Trp Met Lys Arg | g Leu Leu Asp Ile Leu |
|----------------|----------------|-----------------|-----------------------|
| 260            | 265            | 270             |                       |

| Ser Asp | Ile Thr A | Asp Phe Asn (  | Glu Val Val A  | rg Leu Asp Leu Thr Tyr |
|---------|-----------|----------------|----------------|------------------------|
|         | 420       | 425            | 430            |                        |
|         |           |                |                |                        |
| Ile Asp | Asn Trp   | Thr Ile Trp Se | er Asp Ile Lys | Ile Leu Leu Lys Thr    |
| 43      | 5         | 440            | 445            |                        |
|         |           |                |                |                        |
| Val Lys | Val Val   | Leu Leu Arg    | Glu Gly Gly G  | Hn                     |
| 450     |           | 455            |                |                        |
|         |           |                |                |                        |
| <210>   | 17        |                |                |                        |
|         |           |                |                |                        |
| <211>   | 389       |                |                |                        |
|         |           |                |                |                        |
| <212>   | PRT       |                |                |                        |
|         |           |                |                |                        |
| <213>   | Streptoco | occus suis     |                |                        |
|         | •         |                |                |                        |
|         |           |                |                |                        |
| <220>   |           |                |                |                        |

<223> CPS2F

<221> misc\_feature

 $\label{eq:metarg} \mbox{Met Arg Thr Val Tyr Ile Ile Gly Ser Lys Gly Ile Pro Ala Lys Tyr}$ 

1 5 10 15

| Gly Gly Phe G               | ilu Thr Phe V                 | al Glu Lys Leu Thr Glu Tyr Gln Ly          | s Asp |
|-----------------------------|-------------------------------|--|-------|
| 20                          | 25                            | 30   |       |
|                             |                               |  |       |
| Lys Ser Ile As              | n Tyr Phe Va                  | l Ala Cys Thr Arg Glu Asn Ser Ala          | Lys   |
| 35                          | 40                            | 45   |       |
| a                           |                               |  |       |
| · ·                         | •                             | l Phe Glu His Asn Gly Ala Thr Cys          | Phe   |
| 50                          | 55                            | 60   |       |
|                             |                               |  |       |
| Asn Ile Asp V               | al Pro Asn Ile                | e Gly Ser Ala Lys Ala Ile Leu Tyr A        | sp    |
| 65                          | 70                            | 75 80                                      |       |
|                             |                               |  |       |
| Ile Met Ala Le              | u Lys Lys Se                  | r Ile Glu Ile Ala Lys Asp Arg Asn A        | Asp   |
| 85                          | 90                            | 95   |       |
|                             |                               |  |       |
|                             |                               |  |       |
| Thr Ser Pro Ile             | e Phe Tyr Ile l               | Leu Ala Cys Arg Ile Gly Pro Phe Ile        | e     |
| Thr Ser Pro Ile             | e Phe Tyr Ile 1               | Leu Ala Cys Arg Ile Gly Pro Phe Ile        | e     |
|                             |                               |  | e     |
| 100                         | 105                           |  |       |
| 100                         | 105                           | 110  |       |
| 100<br>Tyr Leu Phe L        | 105<br>Lys Lys Gln Ile        | 110<br>e Glu Ser Ile Gly Gly Gln Leu Phe V |       |
| 100<br>Tyr Leu Phe L<br>115 | 105<br>Lys Lys Gln Ile<br>120 | 110<br>e Glu Ser Ile Gly Gly Gln Leu Phe V | √al   |

Arg Gln Tyr Trp Lys Phe Ser Glu Ser Leu Met Leu Lys Tyr Ala Asp

Leu Leu Ile Cys Asp Ser Lys Asn Ile Glu Lys Tyr Ile His Glu Asp

Tyr Arg Lys Tyr Ala Pro Glu Thr Ser Tyr Ile Ala Tyr Gly Thr Asp 180 185 190

Leu Asp Lys Ser Arg Leu Ser Pro Thr Asp Ser Val Val Arg Glu Trp
195 200 205

Tyr Lys Glu Lys Glu Ile Ser Glu Asn Asp Tyr Tyr Leu Val Val Gly
210 215 220

Arg Phe Val Pro Glu Asn Asn Tyr Glu Val Met Ile Arg Glu Phe Met 225 230 235 240

Lys Ser Tyr Ser Arg Lys Asp Phe Val Leu Ile Thr Asn Val Glu His
245 250 255

Asn Ser Phe Tyr Glu Lys Leu Lys Glu Thr Gly Phe Asp Lys Asp
260 265 270

Lys Arg Ile Lys Phe Val Gly Thr Val Tyr Asn Gln Glu Leu Leu Lys
275 280 285

Tyr Ile Arg Glu Asn Ala Phe Ala Tyr Phe His Gly His Glu Val Gly
290 295 300

Gly Thr Asn Pro Ser Leu Leu Glu Ala Leu Ser Ser Thr Lys Leu Asn 305 310 315 320

Leu Leu Asp Val Gly Phe Asn Arg Glu Val Gly Glu Glu Gly Ala 325 330 335

| Lys Tyr Trp | Asn Lys Asp A | Asn Leu His | Arg Val Ile | Asp Ser Cys | Gl |
|-------------|---------------|-------------|-------------|-------------|----|
| 340         | 345           | 34          | 50          |             |    |

Lys Leu Phe Lys Gly

385

<210> 18

<211> 385

<212> PRT

<213> Streptococcus suis

<220>

<221> misc\_feature

<223> CPS2G

<400> 18

| Met Lys Lys Ile Leu Tyr Leu His Ala Gly Ala Glu Leu Tyr Gly Ala |                  |                 |                             |
|---|------------------|-----------------|-----------------------------|
| 1 5   | 10               | 1               | 5                           |
|   |                  |                 |                             |
| Asp Lys Val   | Leu Leu Glu I    | eu Ile Lys G    | ly Leu Asp Lys Asn Glu Phe  |
| 20  | 25               | 30              |                             |
|   |                  |                 |                             |
| Glu Ala His   | Val Ile Leu Pro  | Asn Asp Gl      | y Val Leu Val Pro Ala Leu   |
| 35  | 40               | 45              |                             |
|   |                  |                 |                             |
| Arg Glu Val   | Gly Ala Gln V    | al Glu Val Ile  | Asn Tyr Pro Ile Leu Arg     |
| 50  | 55               | 60              |                             |
|   |                  |                 |                             |
| Arg Lys Tyr   | Phe Asn Pro L    | ys Gly Ile Ph   | e Asp Tyr Phe Ile Ser Tyr   |
| 65  | 70               | 75              | 80                          |
| *** ***   |                  |                 |                             |
| _   | -                | _               | Ala Ile Glu Asn Lys Val     |
| 85  | 90               | 9:              | 5                           |
| A T1 T1 TT  |                  | 701 A1 X7.1     |                             |
| _   |                  |                 | Leu Glu Gly Ile Tyr Leu     |
| 100   | 105              | 110             | J                           |
| Ive Ara Ive   | I an I ve I an D | Pro I au I au T | rp His Val His Glu Ile Ile  |
| 115   | 120              | 125             | ip ins varins ou he he      |
| 113   | 120              | 123             |                             |
| Val Lys Pro Lys Phe Ile Ser Asp Ser Ile Asn Phe Leu Met Gly Arg |                  |                 |                             |
| 130   | 135              | 140             | issi i no Boa wiet diy i ng |
|   |                  | •               |                             |
| Phe Ala Asp Lys Ile Val Thr Val Ser Gln Ala Val Ala Asn His Ile |                  |                 |                             |

Lys Gln Ser Pro His Ile Lys Asp Asp Gln Ile Ser Val Ile Tyr Asn

165 170 175

Gly Val Asp Asn Lys Val Phe Tyr Gln Ser Asp Ala Arg Ser Val Arg

180 185 190

Glu Arg Phe Asp Ile Asp Glu Glu Ala Leu Val Ile Gly Met Val Gly
195 200 205

Arg Val Asn Ala Trp Lys Gly Gln Gly Asp Phe Leu Glu Ala Val Ala 210 215 220

Pro Ile Leu Glu Gln Asn Pro Lys Ala Ile Ala Phe Ile Ala Gly Ser 225 230 235 240

Ala Phe Glu Glu Glu Glu Trp Arg Val Val Glu Leu Glu Lys Lys Ile 245 250 255

Ser Gln Leu Lys Val Ser Ser Gln Val Arg Arg Met Asp Tyr Tyr Ala 260 265 270

Asn Thr Thr Glu Leu Tyr Asn Met Phe Asp Ile Phe Val Leu Pro Ser
275 280 285

Thr Asn Pro Asp Pro Leu Pro Thr Val Val Leu Lys Ala Met Ala Cys 290 295 300

Gly Lys Pro Val Val Gly Tyr Arg His Gly Gly Val Cys Glu Met Val 305 310 315 320 Lys Glu Gly Val Asn Gly Phe Leu Val Thr Pro Asn Ser Pro Leu Asn

325

330

335

Leu Ser Lys Val Ile Leu Gln Leu Ser Glu Asn Ile Asn Leu Arg Lys

340

345

350

Lys Ile Gly Asn Asn Ser Ile Glu Arg Gln Lys Glu His Phe Ser Leu

355

360

365

Lys Ser Tyr Val Lys Asn Phe Ser Lys Val Tyr Thr Ser Leu Lys Val

370

375

380

Tyr

385

<210> 19

<211> 456

<212> PRT

<213> Streptococcus suis

<220>

<221> misc\_feature

<223> cps2h

#### <400> 19

Met Lys Ile Ile Ser Phe Thr Met Val Asn Asn Glu Ser Glu Ile Ile Glu Ser Phe Ile Arg Tyr Asn Tyr Asn Phe Ile Asp Glu Met Val Ile Ile Asp Asn Gly Cys Thr Asp Asn Thr Met Gln Ile Ile Phe Asn Leu Ile Lys Glu Gly Tyr Lys Ile Ser Val Tyr Asp Glu Ser Leu Glu Ala Tyr Asn Gln Tyr Arg Leu Asp Asn Lys Tyr Leu Thr Lys Ile Ile Ala Glu Lys Asn Pro Asp Leu Ile Ile Pro Leu Asp Ala Asp Glu Phe Leu Thr Ala Asp Ser Asn Pro Arg Lys Leu Leu Glu Gln Leu Asp Leu Glu Lys Ile His Tyr Val Asn Trp Gln Trp Phe Val Met Thr Lys Lys Asp 

Asp Ile Asn Asp Ser Phe Ile Pro Arg Arg Met Gln Tyr Cys Phe Glu
130 135 140

Lys Pro Val Trp His His Ser Asp Gly Lys Pro Val Thr Lys Cys Ile Ile Ser Ala Lys Tyr Tyr Lys Lys Met Asn Leu Lys Leu Ser Met Gly His His Thr Val Phe Gly Asn Pro Asn Val Arg Ile Glu His His Asn Asp Leu Lys Phe Ala His Tyr Arg Ala Ile Ser Gln Glu Gln Leu Ile Tyr Lys Thr Ile Cys Tyr Thr Ile Arg Asp Ile Ala Thr Met Glu Asn 

Asn Ile Glu Thr Ala Gln Arg Thr Asn Gln Met Ala Leu Ile Glu Ser 

Gly Val Asp Met Trp Glu Thr Ala Arg Glu Ala Ser Tyr Ser Gly Tyr 

Asp Cys Asn Val Ile His Ala Pro Ile Asp Leu Ser Phe Cys Lys Glu 

Asn Ile Val Ile Lys Tyr Asn Glu Leu Ser Arg Glu Thr Val Ala Glu 

Arg Val Met Lys Thr Gly Arg Glu Met Ala Val Arg Ala Tyr Asn Val 

| Glu Arg Lys Gln Lys Glu Lys Lys Phe Leu Lys Pro Ile Ile Phe Val |                |              |                 |                 |
|---|----------------|--------------|-----------------|-----------------|
| 305   | 310            | 315          | 320             |                 |
|   |                |              |                 |                 |
| Leu Asp G   | ly Leu Lys G   | ly Asp Glu I | Гуг Ile His Pro | Asn Pro Ser Asn |
|   | 325            | 330          | 335             |                 |
|   |                |              |                 |                 |
| His Leu Th  | nr Ile Leu Thr | Glu Met Ty   | r Asn Val Arg   | Gly Leu Leu Thr |
| 340   | 0 3            | 45           | 350             |                 |
|   |                |              |                 |                 |
| Asp Asn H   | is Gln Ile Lys | Phe Leu Ly   | s Val Asn Tyr   | Arg Leu Ile Ile |
| 355   | 360            | 3            | 65              |                 |
|   |                |              |                 |                 |
| Thr Pro As  | sp Phe Ala Ly  | s Phe Leu P  | ro His Glu Phe  | Ile Val Val Pro |
| 370   | 375            | 380          |                 |                 |
|   |                |              |                 |                 |
| Asp Thr Lo  | eu Asp Ile Glu | ı Gln Val Ly | s Ser Gln Tyr   | Val Gly Thr Gly |
| 385   | 390            | 395          | 400             |                 |
|   |                |              |                 |                 |
| Val Asp Leu Ser Lys Ile Ile Ser Leu Lys Glu Tyr Arg Lys Glu Ile |                |              |                 |                 |
|   | 405            | 410          | 415             |                 |
|   | •              |              |                 |                 |
| Gly Phe Ile Gly Asn Leu Tyr Ala Leu Leu Gly Phe Val Pro Asn Met |                |              |                 |                 |
| 420   | •              | 25           | 430             |                 |
|   |                |              |                 |                 |

Leu Asn Arg Ile Tyr Leu Tyr Ile Gln Arg Asn Gly Ile Ala Asn Thr 435 440 445

Ile Ile Lys Ile Lys Ser Arg Leu

|            | _      |       |
|------------|--------|-------|
| < 2.1      | $\sim$ | 2.0   |
| $\sim$ 2.1 | 11/    | - 2.1 |

Met Gln Ala Asp Arg Arg Lys Thr Phe Gly Lys Met Arg Ile Arg Ile

Asn Asn Leu Phe Phe Val Ala Ile Ala Phe Met Gly Ile Ile Ile Ser

 $Asn \ Ser \ Gln \ Val \ Val \ Leu \ Ala \ Ile \ Gly \ Lys \ Ala \ Ser \ Val \ Ile \ Gln \ Tyr$ 

Leu Ser Tyr Leu Val Leu Ile Leu Cys Ile Val Asn Asp Leu Leu Lys

| Asn Asn Lys I  | His Ile Val  | √al Tyr L  | ys Leu C  | Gly Tyr Leu Phe Leu Ile  |
|----------------|--------------|------------|-----------|--------------------------|
| 65             | 70           | 75         | ;         | 80                       |
|                |              |            |           |                          |
| Ile Phe Leu Ph | ne Thr Ile G | ly Ile Cy  | s Gln Glı | n Ile Leu Pro Ile Thr    |
| 85             |              | 90         | 95        |                          |
|                | -            |            | , ,       |                          |
| The Lye He Tu  | r I au Sar I | la Sar M   | et Met Il | o Ilo Sor Vol I ou Alo   |
|                |              |            |           | e Ile Ser Val Leu Ala    |
| 100            | 10:          | 5          | 110       |                          |
|                |              |            |           |                          |
| Thr Leu Pro II | le Ser Leu I | le Lys As  | sp Ile As | p Asp Phe Arg Arg Ile    |
| 115            | 120          |            | 125       |                          |
|                |              |            |           |                          |
| Ser Asn His L  | eu Leu Phe   | Ala Leu    | Phe Ile 7 | Thr Ser Ile Leu Gly Ile  |
| 130            | 135          | 14         |           | in ser ne bea siy ne     |
| 130            | 133          | 19         | 10        |                          |
|                |              |            | ~         |                          |
| Lys Met Gly A  | Ala Thr Met  | Phe Thr    | Gly Ala   | Val Glu Gly Ile Gly Phe  |
| 145            | 150          | 155        |           | 160                      |
|                |              |            |           |                          |
| Ser Gln Gly Pl | he Asn Gly   | Gly Leu    | Thr His 1 | Lys Asn Phe Phe Gly Ile  |
| 165            |              | 170        | 17:       | 5                        |
|                |              |            |           |                          |
| The Ho Lou M   | ot Clu Dho   | Val I au ' | The Tue I | On Alo Tem Less Tem Cler |
|                |              |            | -         | Leu Ala Tyr Lys Tyr Gly  |
| 180            | 18:          | 5          | 190       |                          |
|                |              |            |           |                          |
| Ser Tyr Lys A  | rg Thr Asp   | Arg Phe    | Ile Leu ( | Gly Leu Glu Leu Phe Leu  |
| 195            | 200          |            | 205       |                          |

lle Leu <br/> lle Ser Asn Thr Arg Ser Val Tyr Leu <br/>lle Leu Leu Leu Phe

Leu Phe Leu Val Asn Leu Asp Lys Ile Lys Ile Glu Gln Arg Gln Trp 

Ser Thr Leu Lys Tyr Ile Ser Met Leu Phe Cys Ala Ile Phe Leu Tyr 

Tyr Phe Phe Gly Phe Leu Ile Thr His Ser Asp Ser Tyr Ala His Arg 

Val Asn Gly Leu Ile Asn Phe Phe Glu Tyr Tyr Arg Asn Asp Trp Phe 

His Leu Met Phe Gly Ala Ala Asp Leu Ala Tyr Gly Asp Leu Thr Leu 

Asp Tyr Ala Ile Arg Val Arg Arg Val Leu Gly Trp Asn Gly Thr Leu 

Glu Met Pro Leu Leu Ser Ile Met Leu Lys Asn Gly Phe Ile Gly Leu 

Val Gly Tyr Gly Ile Val Leu Tyr Lys Leu Tyr Arg Asn Val Arg Ile 

Leu Lys Thr Asp Asn Ile Lys Thr Ile Gly Lys Ser Val Phe Ile Ile 

Val Val Leu Ser Ala Thr Val Glu Asn Tyr Ile Val Asn Leu Ser Phe 

| Val Phe Met Pro Ile Cys Phe Cys Leu Leu Asn Ser Ile Ser Thr Met |                  |               |     |  |  |
|---|------------------|---------------|-----|--|--|
| 385   | 390              | 395           | 400 |  |  |
| Glu Ser   | Thr Ile Asn Lys  | Gln Leu Gln 7 | Γhr |  |  |
| <210>   | 21               |               |     |  |  |
| <211>   | 332              |               |     |  |  |
| <212>   | PRT              |               |     |  |  |
| <213>   | Streptococcus si | uis           |     |  |  |
| <220>   |                  |               |     |  |  |

1 5 10 15

<221> misc\_feature

<223> CPS2J

<400> 21

Leu Arg Glu Cys Leu Asp Ser Ile Ile Ser Gln Ser Tyr Thr Asn Leu 20 25 30

Glu Ile Leu Leu Ile Asp Asp Gly Ser Ser Asp Ser Ser Thr Asp Ile Cys Leu Glu Tyr Ala Glu Gln Asp Gly Arg Ile Lys Leu Phe Arg Leu Pro Asn Gly Gly Val Ser Asn Ala Arg Asn Tyr Gly Ile Lys Asn Ser Thr Ala Asn Tyr Ile Met Phe Val Asp Ser Asp Asp Ile Val Asp Gly Asn Ile Val Glu Ser Leu Tyr Thr Cys Leu Lys Glu Asn Asp Ser Asp Leu Ser Gly Gly Leu Leu Ala Thr Phe Asp Gly Asn Tyr Gln Glu Ser Glu Leu Gln Lys Cys Gln Ile Asp Leu Glu Glu Ile Lys Glu Val Arg Asp Leu Gly Asn Glu Asn Phe Pro Asn His Tyr Met Ser Gly Ile Phe 

Phe Asp Thr Glu Gln Trp Leu Gly Glu Asp Leu Leu Phe Asn Leu Asn 180 185 190

Asn Ser Pro Cys Cys Lys Leu Tyr Lys Asn Ile Tyr Ile Asn Gln Gly

Tyr Leu Lys Asn Ile Lys Lys Val Arg Tyr Val Asn Arg Asn Leu Tyr 195 200 205

Phe Ala Arg Arg Ser Leu Gln Ser Thr Thr Asn Thr Phe Lys Tyr Asp

210 215 220

Val Phe Ile Gln Leu Glu Asn Leu Glu Glu Lys Thr Phe Asp Leu Phe 225 230 235 240

Val Lys Ile Phe Gly Gly Gln Tyr Glu Phe Ser Val Phe Lys Glu Thr
245 250 255

Leu Gln Trp His Ile Ile Tyr Tyr Ser Leu Leu Met Phe Lys Asn Gly
260 265 270

Asp Glu Ser Leu Pro Lys Lys Leu His Ile Phe Lys Tyr Leu Tyr Asn 275 280 285

Arg His Ser Leu Asp Thr Leu Ser Ile Lys Arg Thr Ser Ser Val Phe 290 295 300

Lys Arg Ile Cys Lys Leu Ile Val Ala Asn Asn Leu Phe Lys Ile Phe 305 310 315 320

Leu Asn Thr Leu Ile Arg Glu Glu Lys Asn Asn Asp 325 330

<210> 22

- <211> 332
- <212> PRT
- <213> Streptococcus suis
- <220>
- <221> misc\_feature
- <223> CPS2K
- <400> 22

Met Ile Asn Ile Ser Ile Ile Val Pro Ile Tyr Asn Val Glu Gln Tyr

1 5 10 15

Leu Ser Lys Cys Ile Asn Ser Ile Val Asn Gln Thr Tyr Lys His Ile

20 25 30

Glu Ile Leu Leu Val Asn Asp Gly Ser Thr Asp Asn Ser Glu Glu Ile

35 40 45

Cys Leu Ala Tyr Ala Lys Lys Asp Ser Arg Ile Arg Tyr Phe Lys Lys
50 55 60

Glu Asn Gly Gly Leu Ser Asp Ala Arg Asn Tyr Gly Ile Ser Arg Ala
65 70 75 80

Lys Gly Asp Tyr Leu Ala Phe Ile Asp Ser Asp Asp Phe Ile His Ser 85 90 95

Glu Phe Ile Gln Arg Leu His Glu Ala Ile Glu Arg Glu Asn Ala Leu 100 105 110

Val Ala Val Ala Gly Tyr Asp Arg Val Asp Ala Ser Gly His Phe Leu
115 120 125

Thr Ala Glu Pro Leu Pro Thr Asn Gln Ala Val Leu Ser Gly Arg Asn 130 135 140

Val Cys Lys Leu Leu Glu Ala Asp Gly His Arg Phe Val Val Ala 145 150 155 160

Trp Asn Lys Leu Tyr Lys Lys Glu Leu Phe Asp Phe Arg Phe Glu Lys
165 170 175

Gly Lys Ile His Glu Asp Glu Tyr Phe Thr Tyr Arg Leu Leu Tyr Glu
180 185 190

Leu Glu Lys Val Ala Ile Val Lys Glu Cys Leu Tyr Tyr Val Asp 195 200 205

Arg Glu Asn Ser Ile Ile Thr Ser Ser Met Thr Asp His Arg Phe His 210 215 220

Cys Leu Leu Glu Phe Gln Asn Glu Arg Met Asp Phe Tyr Glu Ser Arg
225 230 235 240

| Gly Asp Lys Glu Le | u Leu Leu Glu | ı Cys Tyr Arg Ser | Phe Leu Ala Phe |
|--------------------|---------------|-------------------|-----------------|
| 245                | 250           | 255               |                 |

<213> Streptococcus suis

<220>

<221> misc\_feature

| -000  | ODGOO |
|-------|-------|
| <774> | CPS2O |
|       |       |

# Ser Thr Leu Met Val Ser Ser Ile Ala Phe Phe Leu Pro Ile Phe Gly

| Leu Ser Phe Leu Le | eu Ser Gln | Pro Leu Ser Leu Leu Phe Gly Leu Pr | ю |
|--------------------|------------|------------------------------------|---|
| 100                | 105        | 110                                |   |

| Gln Ile Val P   | he Ser Ser Le  | u Asn Thr V  | al Trp Cys P  | ro Trp Tyr Phe  |
|-----------------|----------------|--------------|---------------|-----------------|
| 260             | 265            | 2            | 70            |                 |
|                 |                |              |               |                 |
| Glu Lys Lys     | Arg Gly Ala A  | Asp Lys Asp  | Leu Leu Ser   | Tyr Val Arg Tyr |
| 275             | 280            | 285          |               |                 |
| True I ou Alo l | lla Chu Lau Dh | o Val The D  | ha Chu Dha I  | on The He Ton   |
| -               | -              |              | ne Giy Phe L  | eu Thr Ile Tyr  |
| 290             | 295            | 300          |               |                 |
| Pro Arg Leu     | Ala Met Leu l  | Leu Glv Glv  | Ser Glu Tvr   | Arg Phe Ser Met |
| 305             | 310            | 315          | 320           |                 |
| 303             | 310            | 313          | 320           |                 |
| Gly Phe Ile P   | ro Met Ile Ile | Val Gly Val  | Phe Phe Val   | Phe Leu Tyr     |
| 32:             |                | 30           | 335           | <b>,</b> -      |
|                 |                |              |               |                 |
| Ser Phe Pro     | Ala Asn Ile Gl | n Phe Tyr S  | er Gly Asn T  | hr Lys Phe Leu  |
| 340             | 345            | 3            | 50            |                 |
|                 |                |              |               |                 |
| Pro Ile Gly T   | hr Phe Ile Ala | Gly Val Le   | u Asn Ile Ser | Val His Phe     |
| 355             | 360            | 365          |               |                 |
|                 |                |              |               |                 |
| Val Leu Ile P   | ro Thr Lys As  | sn Leu Trp ( | Cys Cys Phe   | Ala Thr Thr Ala |
| 370             | 375            | 380          |               |                 |
|                 |                |              |               |                 |
| Ser Tyr Leu I   | Leu Leu Leu V  | Val Leu His  | Tyr Phe Val   | Ala Lys Lys Lys |

Tyr Ala Tyr Asp Glu Val Ala Ile Ser Thr Phe Val Lys Val Ile Ala 405 410 415

Leu Val Val Val Tyr Thr Gly Leu Met Thr Val Phe Val Gly Ser Ile

420

425

430

Trp Ile Arg Trp Ser Leu Gly Ile Ala Val Leu Val Val Tyr Ala Ile

435

440

445

Tyr Phe Arg Lys Glu Leu Thr Val Ala Leu Asn Thr Phe Arg Glu Lys

450

455

460

Arg Ser Lys

465

<210> 24

<211> 338

<212> PRT

<213> Streptococcus suis

<220>

<221> misc\_feature

<223> CPS2P

<400> 24

| Met Val Ty   | r Ile Ile Ala Glu | Ile Gly Cys    | Asn His Asn Gly Asp Val        |
|--------------|-------------------|----------------|--------------------------------|
| 1 5          | 5 10              | 1              | 15                             |
|              |                   |                |                                |
| His Leu Ala  | Arg Lys Met V     | /al Glu Val A  | Ala Val Asp Cys Gly Val Asp    |
| 20           | 25                | 30             |                                |
|              |                   |                |                                |
| Ala Val Lys  | Phe Gln Thr G     | lu Lys Ala A   | sp Leu Leu Ile Ser Lys Tyr     |
| 35           | 40                | 45             |                                |
|              |                   |                |                                |
|              | -                 | -              | r Thr Gly Glu Ser Asp Ser      |
| 50           | 55                | 60             |                                |
| Cla I au Ch  | . Not The Area    | A == I == Cl-  | I an Can Dha Chi Cha Tan I an  |
|              | _                 | _              | Leu Ser Phe Glu Glu Tyr Leu    |
| 65           | 70                | 75             | 80                             |
| A T A        | . A T C 1         |                |                                |
|              |                   | -              | Gly Val Asp Val Phe Ser Thr    |
| 8.           | 5 90              | ç              | 95                             |
| Des Clas Ass | . Ch. Ch. San I   | a A a Dha 1    | Care Ha Care The Area Mad Dura |
| ·            |                   | -              | Leu Ile Ser Thr Asp Met Pro    |
| 100          | 105               | 11             | 0                              |
| Vol Tyr Lug  | Ilo Dro Sor Ch    | Clu Ilo The    | Asn Leu Pro Tyr Leu Glu        |
| •            | •                 |                | Asii Leu Fio Tyl Leu Giu       |
| 115          | 120               | 125            |                                |
| Lvs Ile Glv  | Aro Gln Ala I v   | s I vs Val Ile | Leu Ser Thr Gly Met Ala        |
| 130          | 135               | 140            | Dod Doi 1111 Gry 1viol 1111    |
| 150          | 133               | 140            |                                |
| Val Met Ası  | p Glu Ile His Gl  | n Ala Val Lv   | s Ile Leu Gln Glu Asn Gly      |

| Thr Thr Asp I         |                        | •                  | nr Thr Glu Tyr Pro Tl        | hr Pro    |
|-----------------------|------------------------|--------------------|------------------------------|-----------|
| Tyr Pro Ala L<br>180  | eu Asn Leu A<br>185    |                    | His Thr Leu Lys Lys<br>90    | s Glu Phe |
| Pro Asn Leu 7         | Γhr Ile Gly Tyı<br>200 | Ser Asp H<br>205   | Iis Ser Val Gly Ser G        | ilu Val   |
| Pro Ile Ala Al<br>210 | a Ala Ala Met<br>215   | Gly Ala Gli<br>220 | lu Leu Ile Glu Lys Hi        | is Phe    |
| Thr Leu Asp           | Asn Glu Met C          | ilu Gly Pro        | Asp His Lys Ala Sei          | r Ala Thr |
| 225                   | 230                    | 235                | 240                          |           |
| Pro Asp Ile Le        |                        | •                  | Gly Val Arg Ile Val G<br>255 | lu Gln    |
| Ser Leu Gly L         | ys Phe Glu Ly          | s Glu Pro C        | Glu Glu Val Glu Val          | Arg Asn   |
| 260                   | 265                    | 27                 | 70                           |           |
| Lys Ile Val Al<br>275 | a Glu Lys Ser<br>280   | Ile Val Ala<br>285 | Lys Lys Ala Ile Ala          | Lys       |

Gly Glu Val Phe Thr Glu Glu Asn Ile Thr Val Lys Arg Pro Gly Asn

Gly Ile Ser Pro Met Glu Trp Tyr Lys Val Leu Gly Gln Val Ser Glu

- 325
- 330
- 335

Gln Met

- <210> 25
- <211> 170
- <212> PRT
- <213> Streptococcus suis
- <220>
- <221> misc\_feature
- <223> CPS2Q
- <400> 25

Met Lys Lys Ile Cys Phe Val Thr Gly Ser Arg Ala Glu Tyr Gly Ile

- 1
- 5
- 10
- 15

Met Arg Arg Leu Leu Ser Tyr Leu Gln Asp Asp Pro Glu Met Glu Leu

- 20
- 25
- 30

Asp Leu Val Val Ala Thr Met His Leu Glu Glu Lys Tyr Gly Met Thr

- 35
- 40
- 45

| Val Lys Asp | Ile Glu Ala | Asp Lys Arg Arg | Ile Val Lys Arg Ile Pro |
|-------------|-------------|-----------------|-------------------------|
| 50          | 55          | 60              |                         |

| <213>  | Streptococcus s               | uis           |                             |
|--------|-------------------------------|---------------|-----------------------------|
| <220>  |                               |               |                             |
| <221>  | misc_feature                  |               |                             |
| <223>  | CPS2R                         |               | ·                           |
| <400>  | 26                            |               |                             |
| Met Gl | u Leu Gly Ile As <sub>i</sub> | p Phe Ala Glu | Asp Tyr Tyr Val Val Leu Phe |
| 1      | 5                             | 10            | 15                          |

His Pro Val Thr Leu Glu Asp Asn Thr Ala Glu Glu Gln Thr Gln Ala
20 25 30

Leu Leu Asp Ala Leu Lys Glu Asp Gly Ser Gln Cys Leu Ile Ile Gly
35 40 45

Ser Asn Ser Asp Thr His Ala Asp Lys Ile Met Glu Leu Met His Glu
50 55 60

Phe Val Lys Gln Asp Ser Asp Ser Tyr Ile Phe Thr Ser Leu Pro Thr 65 70 75 80

Arg Tyr Tyr His Ser Leu Val Lys His Ser Gln Gly Leu Ile Gly Asn 85 90 95

| Ser Ser Ser Gly | Leu Ile Glu | Val Pro S | Ser Leu | Gln | Val Pro | Thr | Leu |
|-----------------|-------------|-----------|---------|-----|---------|-----|-----|
| 100             | 105         |           | 110     |     |         |     |     |

Asn Ile Gly Asn Arg Gln Phe Gly Arg Leu Ser Gly Pro Ser Val Val
115 120 125

His Val Gly Thr Ser Lys Glu Ala Ile Val Gly Gly Leu Gly Gln Leu 130 135 140

Arg Asp Val Ile Asp Phe Thr Asn Pro Phe Glu Gln Pro Asp Ser Ala
145 150 155 160

Leu Gln Gly Tyr Arg Ala Ile Lys Glu Phe Leu Ser Val Gln Ala Ser 165 170 175

Thr Met Lys Glu Phe Tyr Asp Arg 180

<210> 27

<211> 208

<212> PRT

<213> Streptococcus suis

<220>

<221> misc\_feature

#### <223> CPS2S

<400> 27

Met Lys Lys Val Ala Phe Leu Gly Ala Gly Thr Phe Ser Asp Gly Val

1 5 10 15

Leu Pro Trp Leu Asp Arg Thr Arg Tyr Glu Leu Ile Gly Tyr Phe Glu
20 25 30

Asp Lys Pro Ile Ser Asp Tyr Arg Gly Tyr Pro Val Phe Gly Pro Leu

35 40 45

Gln Asp Val Leu Thr Tyr Leu Asp Asp Gly Lys Val Asp Ala Val Phe
50 55 60

Val Thr Ile Gly Asp Asn Val Lys Arg Lys Glu Ile Phe Asp Leu Leu 65 70 75 80

Ala Lys Asp His Tyr Asp Ala Leu Phe Asn Ile Ile Ser Glu Gln Ala 85 90 95

Asn Ile Phe Ser Pro Asp Ser Ile Lys Gly Arg Gly Val Phe Ile Gly
100 105 110

Phe Ser Ser Phe Val Gly Ala Asp Ser Tyr Val Tyr Asp Asn Cys Ile
115 120 125

Ile Asn Thr Gly Ala Ile Val Glu His His Thr Thr Val Glu Ala His

130 135 140

 $Cys\ Asn\ Ile\ Thr\ Pro\ Gly\ Val\ Thr\ Ile\ Asn\ Gly\ Leu\ Cys\ Arg\ Ile\ Gly$ 

- 145
- 150
- 155

160

Glu Ser Thr Tyr Ile Gly Ser Gly Ser Thr Val Ile Gl<br/>n Cys Ile Glu

- 165
- 170
- 175

Ile Ala Pro Tyr Th<br/>r Thr Leu Gly Ala Gly Thr Val Val Leu Lys Ser  $\,$ 

- 180
- 185
- 190

Leu Thr Glu Ser Gly Thr Tyr Val Gly Val Pro Ala Arg Lys Ile Lys

- 195
- 200
- 205

<210> 28

- <211> 410
- <212> PRT
- <213> Streptococcus suis
- <220>
- <221> misc\_feature
- <223> CPS2T
- <400> 28

| Met Glu Pro I  | le Cys Leu Ile | Pro Ala Ar   | g Ser Gly Ser Lys Gly Leu   |
|----------------|----------------|--------------|-----------------------------|
| 1 5            | 10             |              | 15                          |
|                |                |              |                             |
| Pro Asn Lys A  | Asn Met Leu F  | he Leu Asp   | Gly Val Pro Met Ile Phe His |
| 20             | 25             | 30           |                             |
|                |                |              |                             |
| Thr Ile Arg Al | la Ala Ile Glu | Ser Gly Cys  | Phe Lys Lys Glu Asn Ile     |
| 35             | 40             | 45           |                             |
|                |                |              |                             |
| Tyr Val Ser T  | hr Asp Ser Gl  | u Val Tyr L  | ys Glu Ile Cys Glu Thr Thr  |
| 50             | 55             | 60           |                             |
|                |                |              |                             |
| Gly Val Gln V  | al Leu Met A   | rg Pro Ala   | Asp Leu Ala Thr Asp Phe Thr |
| 65             | 70             | 75           | 80                          |
|                |                |              | •                           |
| Thr Ser Phe G  | ln Leu Asn G   | lu His Phe I | Leu Gln Asp Phe Ser Asp Asp |
| 85             | 90             | ģ            | 95                          |
|                |                |              |                             |
| Gln Val Phe V  | al Leu Leu G   | ln Val Thr S | Ser Pro Leu Arg Ser Gly Lys |
| 100            | 105            | 11           | 0                           |
|                |                |              |                             |
| His Val Lys G  | lu Ala Met Gl  | u Leu Tyr (  | Gly Lys Gly Gln Ala Asp His |
| 115            | 120            | 125          |                             |
|                |                |              |                             |
| Val Val Ser Pl | ne Thr Lys Va  | l Asp Lys S  | er Pro Thr Leu Phe Ser Thr  |
| 130            | 135            | 140          |                             |
|                |                |              |                             |
| Leu Asp Glu A  | Asn Gly Phe A  | la Lys Asp   | Ile Ala Gly Leu Gly Gly Ser |

Tyr Arg Arg Gln Asp Glu Lys Thr Leu Tyr Tyr Pro Asn Gly Ala Ile 165 170 175

Tyr Ile Ser Ser Lys Gln Ala Tyr Leu Ala Asp Lys Thr Tyr Phe Ser 180 185 190

Glu Lys Thr Ala Ala Tyr Val Met Thr Lys Glu Asp Ser Ile Asp Val 195 200 205

Asp Asp His Phe Asp Phe Thr Gly Val Ile Gly Arg Ile Tyr Phe Asp 210 215 220

Tyr Gln Arg Arg Glu Gln Gln Asn Lys Pro Phe Tyr Lys Arg Glu Leu 225 230 235 240

Lys Arg Leu Cys Glu Gln Arg Val His Asp Ser Leu Val Ile Gly Asp 245 250 255

Ser Arg Leu Leu Ala Leu Leu Asp Gly Phe Asp Asn Ile Ser Ile
260 265 270

Gly Gly Met Thr Ala Ser Thr Ser Leu Glu Asn Gln Gly Leu Phe Leu 275 280 285

Ala Thr Pro Ile Lys Lys Val Leu Leu Ser Leu Gly Val Asn Asp Leu 290 295 300

Ile Thr Asp Tyr Pro Leu His Met Ile Glu Asp Thr Ile Arg Gln Leu 305 310 315 320

| Met | Glu Se | r Leu | Val Sei | Lys | Ala Glu | ı Gln | Val Glu | Val | Thr | Thr | Ile |
|-----|--------|-------|---------|-----|---------|-------|---------|-----|-----|-----|-----|
|     |        |       |         |     |         |       |         |     |     |     |     |

325

330

335

Ala Tyr Thr Leu Phe Arg Asp Ser Val Ser Asn Glu Glu Thr Val Gln
340 345 350

## Asn Gln Leu Ile Leu Thr Ser Leu Thr Arg

405

410

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<211> 6992

<212> DNA

<213> Streptococcus suis

<220>

<221> misc\_feature

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| <210> 3 | 30 |
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|---------|----|

Ala Ile Leu Thr Ser His Ile Pro Asn Ala Asp Leu Asn Arg Ser Gly

Ile Phe Ile Ile Met Met Val His Tyr Phe Ala Phe Phe Ile Ser Arg

Met Pro Val Glu Phe Glu Tyr Arg Gly Asn Leu Ile Glu Phe Glu Lys

Thr Phe Asn Tyr Ser Ile Ile Phe Ala Ile Phe Leu Thr Ala Val Ser Phe Leu Leu Glu Asn Asn Phe Ala Leu Ser Arg Arg Gly Ala Val Tyr Phe Thr Leu Ile Asn Phe Val Leu Val Tyr Leu Phe Asn Val Ile Ile Lys Gln Phe Lys Asp Ser Phe Leu Phe Ser Thr Ile Tyr Gln Lys Lys Thr Ile Leu Ile Thr Thr Ala Glu Arg Trp Glu Asn Met Gln Val Leu Phe Glu Ser His Lys Gln Ile Gln Lys Asn Leu Val Ala Leu Val Val Leu Gly Thr Glu Ile Asp Lys Ile Asn Leu Ser Leu Pro Leu Tyr Tyr 

Ser Val Glu Glu Ala Ile Glu Phe Ser Thr Arg Glu Val Val Asp His 180 185 190

Val Phe Ile Asn Leu Pro Ser Glu Phe Leu Asp Val Lys Gln Phe Val 195 200 205

Ser Asp Phe Glu Leu Leu Gly Ile Asp Val Ser Val Asp Ile Asn Ser 210 215 220

| Phe Gly   | Phe Thr Ala Le | u Lys Asn Lys  | Lys Ile Gln Leu L | eu Gly Asp |
|-----------|----------------|----------------|-------------------|------------|
| 225       | 230            | 235            | 240               |            |
| His Ser I | le Val Thr Phe | Ser Thr Asn Ph | e Tvr Lvs Pro Se  | r His Ile  |

| Phe Glu          | Lys Tyr T   | hr Pro Gly | Gln Lys    | Arg Arg Leu   | Ser Phe Lys Pro |  |
|------------------|-------------|------------|------------|---------------|-----------------|--|
| 385              | 390         |            | 395        | 400           |                 |  |
|                  |             |            |            |               |                 |  |
| Gly Ile 7        | Thr Gly Le  | u Trp Gln  | Val Ser G  | ly Arg Ser A  | sn Ile Thr Asp  |  |
|                  | 405         | 410        |            | 415           |                 |  |
|                  |             |            |            |               |                 |  |
| Phe Asp          | Asp Val V   | /al Arg Le | eu Asp Lei | ı Ala Tyr Ile | Asp Asn Trp Thr |  |
| 4                | 420         | 425        | 4          | 30            |                 |  |
|                  |             |            |            |               |                 |  |
| Ile Trp S        | Ser Asp Ile | Lys Ile Le | eu Leu Ly  | s Thr Val Lys | s Val Val Leu   |  |
| 435              | 5           | 440        | 445        |               |                 |  |
|                  |             |            |            |               |                 |  |
| Leu Arg          | Glu Gly S   | er Lys     |            |               |                 |  |
| 450              |             |            |            |               |                 |  |
|                  |             |            |            |               |                 |  |
| <210> 3          | 31          |            |            |               |                 |  |
|                  |             |            |            |               |                 |  |
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| <212> 1          | DD T        |            |            |               |                 |  |
| <212> 1          | PKI         |            |            |               |                 |  |
| <213> 9          | Streptococ  | cue enie   |            |               |                 |  |
| ~21 <i>J</i> ~ \ | энсриссос   | cus suis   |            |               |                 |  |
| <220>            |             |            |            |               |                 |  |

<221> misc\_feature

<223> CPS1F

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#### Leu Gly Ser Ile Phe

### 

Met Ile Phe Val Thr Val Gly Thr His Glu Gln Gln Phe Asn Arg Leu

Ile Lys Glu Ile Asp Leu Leu Lys Lys Asn Gly Ser Ile Thr Asp Glu

Ile Phe Ile Gln Thr Gly Tyr Ser Asp Tyr Ile Pro Glu Tyr Cys Lys

| Tyr Lys Lys | Phe Leu Ser | Tyr Lys Glu Met Glu | ı Gln Tyr Ile Asn Lys |
|-------------|-------------|---------------------|-----------------------|
| 50          | 55          | 60                  |                       |

Gln Glu Asn Glu

| <213>  | Streptococcus suis  |
|--------|---|
| <220>  |   |
| <221>  | misc_feature  |
| <223>  | CPS1H   |
| <400>  | 33 .  |
| Met Ph | e Lys Leu Phe Lys Tyr Asp Pro Glu Tyr Phe Ile Phe Lys Tyr |

10

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5

Phe Trp Leu Ile Ile Phe Ile Pro Glu Gln Lys Tyr Val Phe Leu Leu 20 25 30

15

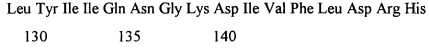
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Leu Ile Leu Lys Asn Glu Ile Leu Leu Phe Leu Leu Trp Ser Ile Leu 50 55 60

Cys Phe Val Ser Val Val Thr Ser Met Phe Val Glu Ile Asn Phe Glu
65 70 75 80

Arg Leu Phe Ala Asp Phe Thr Ala Pro Ile Ile Trp Ile Ile Ala Ile
85 90 95

| Met Tyr Tyr As  | sn Leu Tyr Ser | Phe Ile Asn Ile As | p Tyr Lys Lys Le   |
|-----------------|----------------|--------------------|--------------------|
| 100             | 105            | 110                |                    |
| Lys Asn Ser Ile | Phe Phe Ser P  | he Leu Val Leu Le  | eu Gly Ile Ser Ala |
| 115             | 120            | 125                |                    |
| I T II. II.     | Cla A a Claste | A II . X/-1 Dl     | T A A TT'-         |



| Ala Val Tyr | Asn Ser Arg Glu | Ser Ser Asn Glu A | Ala Arg Phe Ile Ile |
|-------------|-----------------|-------------------|---------------------|
| 260         | 265             | 270               |                     |

Tyr Gln Gly Ser Ile Asp Lys Val Leu Glu Asn Asn Ile Leu Phe Gly 

Tyr Gly Ile Ser Glu Tyr Ser Val Thr Gly Thr Trp Leu Gly Ser His 

Ser Gly Tyr Ile Ser Phe Phe Tyr Lys Ser Gly Ile Val Gly Leu Ile 

Leu Leu Met Phe Ser Phe Phe Tyr Val Ile Lys Lys Ser Tyr Gly Val 

Asn Gly Glu Thr Ala Leu Phe Tyr Phe Thr Ser Leu Ala Ile Phe Phe 

Ile Tyr Glu Thr Ile Asp Pro Ile Ile Ile Ile Leu Val Leu Phe Phe 

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Thr Lys Asn Glu

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- <213> Streptococcus suis
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- <221> misc\_feature
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- Tyr Leu Asp Lys Cys Ile Asn Ser Ile Ile Asn Gln Thr Tyr Thr Asn
  - 20

- 25
- 30
- Leu Glu Val Ile Leu Val Asn Asp Gly Ser Thr Asp Asp Ser Glu Lys
  - 35
- 40
- 45
- Ile Cys Leu Asn Tyr Met Lys Asn Asp Gly Arg Ile Lys Tyr Tyr Lys
  - 50
- 55
- 60
- Lys Ile Asn Gly Gly Leu Ala Asp Ala Arg Asn Phe Gly Leu Glu His
- 65
- 70
- 75
- 80

| Ala Thr Gly Lys | Tyr Ile Ala Phe | Val Asp Ser | Asp Asp Ty | r Ile Glu |
|-----------------|-----------------|-------------|------------|-----------|
| 85              | 90              | 95          |            |           |

| Glu Phe Ser | His Tyr I | Phe Asp | Ala Lys | Val Ile L | ys Glu Lys | Val Lys |
|-------------|-----------|---------|---------|-----------|------------|---------|
|             |           |         |         |           |            |         |

Lys Gln

<220>

<223> CPS1J

<400> 35

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1 5 10 15

Leu Ser Ser Cys Ile Glu Ser Ile Ile Asn Gln Asn Tyr Lys Asn Ile

20 25 30

Glu Ile Leu Leu Ile Asp Asp Gly Ser Val Asp Asp Ser Ala Lys Ile

35 40 45

Cys Lys Glu Tyr Glu Lys Asp Lys Arg Val Lys Ile Phe Phe Thr Asn

50 55 60

His Ser Gly Val Ser Asn Ala Arg Asn His Gly Ile Lys Arg Ser Thr

65 70 75 80

Ala Glu Tyr Ile Met Phe Val Asp Ser Asp Asp Val Val Asp Ser Arg

85 90 95

Leu Val Glu Lys Leu Tyr Phe Asn Ile Ile Lys Ser Arg Ser Asp Leu

100 105 110

Ser Gly Cys Leu Tyr Ala Thr Phe Ser Glu Asn Ile Asn Asn Phe Glu

115 120 125

| Val Asn | Asn Pro Asn Ile | Asp Phe Glu Ala | Ile Asn Thr Val Gln Asp |
|---------|-----------------|-----------------|-------------------------|
| 130     | 135             | 140             |                         |
|         |                 |                 |                         |

Met Gly Glu Lys Asn Phe Met Asn Leu Tyr Ile Asn Asn Ile Phe Ser

145 150 155 160

Thr Pro Val Cys Lys Leu Tyr Lys Lys Arg Tyr Ile Thr Asp Leu Phe 165 170 175

Gln Glu Asn Gln Trp Leu Gly Glu Asp Leu Leu Phe Asn Leu His Tyr 180 185 190

Leu Lys Asn Ile Asp Arg Val Ser Tyr Leu Thr Glu His Leu Tyr Phe
195 200 205

Tyr Arg Arg Gly Ile Leu Ser Thr Val Asn Ser Phe Lys Glu Gly Val 210 215 220

Phe Leu Gln Leu Glu Asn Leu Gln Lys Gln Val Ile Val Leu Phe Lys 225 230 235 240

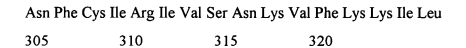
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245 250 255

Arg Trp Gln Val Phe Tyr Tyr Ser Leu Leu Met Phe Lys Tyr Gly Lys 260 265 270

Gln Ser Ile Phe Asp Lys Phe Leu Ile Phe Arg Asn Leu Tyr Lys Lys 275 280 285

| Tyr Tyr Phe | Asn Leu Leu | Lys Val Ser Asn Lys Asn Ser Leu Se | r Lys |
|-------------|-------------|------------------------------------|-------|
| 290         | 295         | 300                                |       |



Trp Leu

Met Asp Thr Ile Ser Lys Ile Ser Ile Ile Val Pro Ile Tyr As<br/>n Val  $\,$ 

1 5 10 15

Glu Lys Tyr Leu Ser Lys Cys Ile Asp Ser Ile Val Asn Gln Thr Tyr Lys His Ile Glu Ile Leu Leu Val Asn Asp Gly Ser Thr Asp Asn Ser Glu Glu Ile Cys Leu Ala Tyr Ala Lys Lys Asp Ser Arg Ile Arg Tyr Phe Lys Lys Glu Asn Gly Gly Leu Ser Asp Ala Arg Asn Tyr Gly Ile Ser Arg Ala Lys Gly Asp Tyr Leu Ala Phe Ile Asp Ser Asp Asp Phe Ile His Ser Glu Phe Ile Gln Arg Leu His Glu Ala Ile Glu Arg Glu Asn Ala Leu Val Ala Val Ala Gly Tyr Asp Arg Val Asp Ala Ser Gly His Phe Leu Thr Ala Glu Pro Leu Pro Thr Asn Gln Ala Val Leu Ser 

Val Val Ala Cys Asn Lys Leu Tyr Lys Lys Glu Leu Phe Glu Asp Phe 165 170 175

Gly Arg Asn Val Cys Lys Leu Leu Glu Ala Asp Gly His Arg Phe

| Arg Phe Glu | Lys Gly Lys Ile | His Glu Asp Glu | u Tyr Phe T | hr Tyr Arg |
|-------------|-----------------|-----------------|-------------|------------|
| 180         | 185             | 190             |             |            |

<213> Streptococcus suis

<220>

<221> misc\_feature

<223> CPS9

<400> 37 60 aagcttatcg tcaaggtgtt cgctatatcg tggcgacatc tcatagacga aaagggatgt ttgaaacacc agaaaaagtt atcatgacta actttcttca atttaaagac gcagtagcag 120 aagtttatcc tgaaatacga ttgtgctatg gtgctgaatt gtattatagt aaagatatat 240 taagcaaact tgaaaaaaag aaagtaccca cacttaatgg ctcgcgctat attcttttgg 300 agttcagtag tgatactcct tggaaagaga ttcaagaagc agtgaacgaa gtgacgctac ttgggctaac tcccgtactt gcccatatag aacgatatga cgccctagcg tttcatgcag 360 agagagtaga agagttaatt gacaagggat gctatactca ggtaaatagt aatcatgtgc 420 tgaagcccac tttaattggt gatcgagcaa aagaatttaa aaaacgtact cggtattttt 480 540 tagagcagga tttagtacat tgtgttgcta gcgatatgca taatttatct agtagacctc cgtttatgag ggaggcttat aagttgctaa cagaggaatt tggcaaagat aaagcgaaag 600

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catgggttgt ctctcaaaat aaatctaatc aggggcatta tcaaacattt ataaatttga 4320
caaagttagt tcaggaagga atagtctttt tttcagatca agatgatatt tgggactgtc 4380
ataaaattga gacaatgctt ccaatctttg acagagaaaa tgtatcaatg gtgttttgca 4440
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tcaatacgta ctctctaga 4519

- <210> 38
- <211> 215
- <212> PRT
- <213> Streptococcus suis
- <220>
- <221> misc\_feature
- <223> CPS9D
- <400> 38

Ala Tyr Arg Gln Gly Val Arg Tyr Ile Val Ala Thr Ser His Arg Arg Lys Gly Met Phe Glu Thr Pro Glu Lys Val Ile Met Thr Asn Phe Leu Gln Phe Lys Asp Ala Val Ala Glu Val Tyr Pro Glu Ile Arg Leu Cys Tyr Gly Ala Glu Leu Tyr Tyr Ser Lys Asp Ile Leu Ser Lys Leu Glu Lys Lys Val Pro Thr Leu Asn Gly Ser Arg Tyr Ile Leu Leu Glu Phe Ser Ser Asp Thr Pro Trp Lys Glu Ile Gln Glu Ala Val Asn Glu Val Thr Leu Leu Gly Leu Thr Pro Val Leu Ala His Ile Glu Arg Tyr Asp Ala Leu Ala Phe His Ala Glu Arg Val Glu Glu Leu Ile Asp Lys Gly Cys Tyr Thr Gln Val Asn Ser Asn His Val Leu Lys Pro Thr Leu Ile Gly Asp Arg Ala Lys Glu Phe Lys Lys Arg Thr Arg Tyr Phe Leu

| Glu G | iln Asp | Leu V | Val His | Cys | Val | Ala | Ser | Asp | Met | His | Asn | Leu | Ser |
|-------|---------|-------|---------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
|-------|---------|-------|---------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|

165

170

175

Ser Arg Pro Pro Phe Met Arg Glu Ala Tyr Lys Leu Leu Thr Glu Glu 180 185 190

Phe Gly Lys Asp Lys Ala Lys Ala Leu Leu Lys Lys Asn Pro Leu Met 195 200 205

Leu Leu Lys Asn Gln Ala Ile

210

215

<210> 39

<211> 608

<212> PRT

<213> Streptococcus suis

<220>

<221> misc\_feature

<223> CPS9E

<400> 39

| Met Asp Leu    | Gly Thr Val T   | Thr Asp Lys Le   | u Leu Glu Arg Asn Ser Lys |
|----------------|-----------------|------------------|---------------------------|
| 1 5            | 10              | 15               |                           |
|                |                 |                  |                           |
| Arg Leu Ile I  | Leu Val Cys M   | et Asp Thr Cys   | Leu Leu Ile Val Ser Met   |
| 20             | 25              | 30               |                           |
|                |                 |                  |                           |
| Ile Leu Ser A  | arg Leu Phe Le  | eu Asp Val Ile I | le Asp Ile Pro Asp Glu    |
| 35             | 40              | 45               |                           |
|                |                 |                  |                           |
| Arg Phe Ile I  | Leu Ala Val Le  | u Phe Val Ser I  | le Leu Tyr Leu Ile Leu    |
| 50             | 55              | 60               |                           |
|                |                 |                  |                           |
| Ser Phe Arg    | Leu Lys Val Pl  | he Ser Leu Ile   | Thr Arg Tyr Thr Gly Tyr   |
| 65             | 70              | 75               | 80                        |
|                |                 |                  |                           |
|                |                 |                  | e Ser Ala His Ser Leu     |
| 85             | 90              | 95               |                           |
|                |                 |                  |                           |
|                |                 | -                | la Phe Ser Tyr Arg Phe    |
| 100            | 105             | 110              |                           |
| Ila I au Wal C | an I an Dha I a | C T \$7-1 \$     | And I am II a Thin Don A  |
|                |                 |                  | Met Leu Ile Thr Pro Arg   |
| 115            | 120             | 125              |                           |
| Ile Val Trn I  | ve Val I au Hie | Clu The Ara I    | Lys Asn Ala Ile Arg Lys   |
| 130            | 135             | 140              | Lys Asii Aid lie Aig Lys  |
| 150            | 133             | 170              |                           |
| Lys Asp Ser    | Pro Leu Arg Il  | e Leu Val Val (  | Gly Ala Gly Asp Gly Gly   |

| Asn Ile Phe Ile Asn | Thr Val Lys Asp | Arg Lys Leu | Asn Phe Glu Ile |
|---------------------|-----------------|-------------|-----------------|
| 165                 | 170             | 175         |                 |

Val Gly Ile Val Asp Arg Asp Pro Asn Lys Leu Gly Thr Phe Ile Arg
180 185 190

Thr Ala Lys Val Leu Gly Asn Arg Asn Asp Ile Pro Arg Leu Val Glu
195 200 205

Glu Leu Ala Val Asp Gln Val Thr Ile Ala Ile Pro Ser Leu Asn Gly
210 215 220

Lys Glu Arg Glu Lys Ile Val Glu Ile Cys Asn Thr Thr Gly Val Thr
225 230 235 240

Val Asn Asn Met Pro Ser Ile Glu Asp Ile Met Ala Gly Asn Met Ser

245 250 255

Val Ser Ala Phe Gln Glu Ile Asp Val Ala Asp Leu Leu Gly Arg Pro 260 265 270

Glu Val Val Leu Asp Gln Asp Glu Leu Asn Gln Phe Phe Gln Gly Lys
275 280 285

Thr Ile Leu Val Thr Gly Ala Gly Gly Ser Ile Gly Ser Glu Leu Cys 290 295 300

Arg Gln Ile Ala Lys Phe Thr Pro Lys Arg Leu Leu Leu Gly His
305 310 315 320

Gly Glu Asn Ser Ile Tyr Leu Ile His Arg Glu Leu Leu Glu Lys Tyr

325 330 335

Gln Gly Lys Ile Glu Leu Val Pro Leu Ile Ala Asp Ile Gln Asp Arg 340 345 350

Glu Leu Ile Phe Ser Ile Met Ala Glu Tyr Gln Pro Asp Val Val Tyr 355 360 365

His Ala Ala His Lys His Val Pro Leu Met Glu Tyr Asn Pro His 370 375 380

Glu Ala Val Lys Asn Asn Ile Phe Gly Thr Lys Asn Val Ala Glu Ala 385 390 395 400

Ala Lys Thr Ala Lys Val Ala Lys Phe Val Met Val Ser Thr Asp Lys
405 410 415

Ala Val Asn Pro Pro Asn Val Met Gly Ala Thr Lys Arg Val Ala Glu 420 425 430

Met Ile Val Thr Gly Leu Asn Glu Pro Gly Gln Thr Gln Phe Ala Ala 435 440 445

Val Arg Phe Gly Asn Val Leu Gly Ser Arg Gly Ser Val Val Pro Leu 450 455 460

Phe Lys Glu Gln Ile Arg Lys Gly Gly Pro Val Thr Val Thr Asp Phe 465 470 475 480

| Arg Met | Thr Arg | Tyr Phe Met | Thr Ile Pro Glu | ı Ala Ser | Arg Leu | Val |
|---------|---------|-------------|-----------------|-----------|---------|-----|
|         | 105     | 400         | 405             |           |         |     |

485 490 495

<210> 40

<211> 200

<212> PRT

| <213>   | Streptod  | coccus suis   |                   |                 |                 |
|---------|-----------|---------------|-------------------|-----------------|-----------------|
| <220>   |           |               |                   |                 |                 |
| <221>   | misc_fea  | ature         |                   |                 |                 |
| <223>   | CPS9F     |               |                   |                 |                 |
| <400>   | 40        |               |                   |                 |                 |
| Met Ty  | r Pro Ile | Cys Lys Arg   | ; Ile Leu Al      | a Ile Ile Ile S | Ser Gly Ile     |
| 1       | 5         | 10            | 1                 | 15              |                 |
| Ala Ile | Val Val 1 | Leu Ser Pro   | Ile Leu Leu       | Leu Ile Ala     | Leu Ala Ile     |
|         | 20        | 25            | 30                |                 |                 |
| Lys Let |           | r Lys Gly Pro | o Val Leu F<br>45 | he Lys Gln      | Lys Arg Val Gly |
| 33      |           | 40            | 43                |                 |                 |
| Lys Ası | ı Lys Sei | r Tyr Phe Me  | t Ile Tyr L       | ys Phe Arg S    | Ser Met Tyr Val |
| 50      |           | 55            | 60                |                 |                 |
|         |           |               |                   |                 |                 |

Met Ile Thr Lys Val Gly Ala Phe Leu Arg Lys Thr Ser Leu Asp Glu 85 90 95

Asp Ala Pro Ser Asp Met Pro Thr His Leu Leu Lys Asp Pro Lys Ala

| Leu Pro Gln Leu | Phe Asn Ile P | he Lys Gly Glu Met | Ala Ile Val Gly |
|-----------------|---------------|--------------------|-----------------|
| 100             | 105           | 110                |                 |

<213> Streptococcus suis

<220>

<221> misc\_feature

<223> CPS2G

<400> 41

Met Lys Phe Ser Val Leu Met Ser Val Tyr Glu Lys Glu Lys Pro Glu

1 5 10 15

Phe Leu Arg Glu Ser Leu Glu Ser Ile Leu Val Asn Gln Thr Met Ile
20 25 30

Pro Thr Glu Val Val Leu Val Glu Asp Gly Pro Leu Asn Gln Ser Leu 35 40 45

Tyr Ser Ile Leu Glu Glu Phe Lys Ser Arg Phe Ser Phe Phe Lys Thr
50 55 60

Ile Ala Leu Glu Lys Asn Ser Gly Leu Gly Ile Ala Leu Asn Glu Gly
65 70 75 80

Leu Lys His Cys Asn Tyr Glu Trp Val Cys Thr Lys Trp Ile Leu Met 85 90 95

Met Leu His Ile His Thr Arg Phe Glu Lys Gln Val Asn Phe Ile Lys
100 105 110

Ser Glu Ile Val Ser His Lys Asn Val Pro Thr Gln His Asp Glu Ile

Leu Lys Met Ala Arg Arg Glu Lys Ser Met Cys His Met Thr Val Met

Phe Lys Lys Ser Val Glu Arg Ala Gly Gly Tyr Gln Thr Leu Pro

Tyr Val Glu Asp Tyr Phe Leu Trp Val Arg Met Ile Ala Ser Gly Ser

Lys Phe Ala Asn Ile Asp Glu Thr Leu Val Leu Ala Arg Val Gly Asn

Gly Met Phe Asn Arg Gly Asn Arg Glu Gln Ile Asn Ser Trp Thr

Leu Leu Ile Glu Phe Met Leu Ala Gln Gly Ile Val Thr Pro Leu Asp

Val Phe Ile Asn Gln Ile Tyr Ile Arg Val Phe Val Tyr Met Pro Thr

Trp Ile Lys Lys Leu Ile Tyr Gly Lys Ile Leu Arg Lys

- <210> 42
- <211> 143
- <212> PRT
- <213> Streptococcus suis
- <220>
- <221> misc\_feature
- <223> CPS9H
- <400> 42

Met Ile Thr Val Leu Met Ala Thr Tyr Asn Gly Ser Pro Phe Ile Ile

- 1
- 15

Lys Gln Leu Asp Ser Ile Arg Asn Gln Ser Val Ser Ala Asp Lys Val

20

5

25

10

30

Ile Ile Trp Asp Asp Cys Ser Thr Asp Asp Thr Ile Lys Ile Ile Lys

- 35
- 40
- 45

 $Asp\ Tyr\ Ile\ Lys\ Lys\ Tyr\ Ser\ Leu\ Asp\ Ser\ Trp\ Val\ Val\ Ser\ Gln\ Asn$ 

- 50
- 55
- 60

| Lys Ser | Asn Gln Gly | His Tyr Gln T | hr Phe Ile Asn | Leu Thr Lys Leu |
|---------|-------------|---------------|----------------|-----------------|
| 65      | 70          | 75            | 80             |                 |

<400> 43 60 ctgcagcaca taagcatgtt ccattgatgg aatataatcc acatgaagca gtgaagaata atatttttgg aacgaagaat gtggctgagg cggctaaaac tgcaaaggtt gccaaatttg 120 180 ttatggtttc aacagataaa gctgttaatc cgccaaatgt catgggagcg actaaacgtg ttgcagaaat gattgtaaca ggtttaaacg agccaggtca gactcaattt gcggcagtcc 240

300 gttttgggaa tgttctaggt agtcgtggaa gtgttgttcc gctattcaaa gagcaaatta 360 gaaaaggtgg acctgttacg gttaccgact ttaggatgac tcgttatttc atgacgattc 420 ctgaggcaag tcgtttggtt atccaagctg gacatttggc aaaaggtgga gaaatctttg tcttggatat gggtgagcca gtacaaatcc tggaattggc aagaaaagtt atcttgttaa 480 gcggacatac agaggaagaa atcgggattg tagaatctgg aatcagacca ggcgagaaac 540 tctacgagga attgttatca acagaagaac gtgtcagcga acagattcat gaaaaaatat 600 660 ttgtgggtcg cgttacaaat aagcagtcgg acattgtcaa ttcatttatc aatggattac 720 aaagtaaaaa atatttttac tttcctagag tttaaacgat gtttaagttc taggaaggtt 780

ggaattgett tegtggaggt gatagataga aacetatata tttgtagaag aaaggatatt 840

aaactaaagg tgaatcggaa cataaagttt agatagagtt ggtatttaat gccaaacagg 900 960 tgaatgcaac ctctcgctcg ttactaagca ggagatagta aagttgcttg aaagagagtt tgttaatcag tataagtagg ctaaagtgag aatatatatc tattattatc ggtaatgata 1020 ctattattga gaattattgt agtggggata aaaataattt ttggtgattt tatcgtccga 1080 cttaaaggtg ggttaaaaaa gtacttatat tcttttagaa ttgatgaaaa atatggggga 1140 atataatatt tataggagat acgatgacta gagtagagtt gattactaga gaatttttta 1200 agaagaatga agcaaccagt aaatattttc agaagataga atcaagaaga ggtgaattat 1260 ttattaaatt ctttatggat aagttacttg cgcttatcct attattgcta ttatccccag 1320 taatcattat attagctatt tggataaaat tagatagtaa ggggccaatt ttttatcgcc 1380 aagaacgtgt tacgagatat ggtcgaattt ttagaatatt taagtttaga acaatgattt 1440 ctgatgcgga taaagtcgga agtcttgtca cagtcggtca agataatcgt attacgaaag 1500 tcggtcacat tatcagaaaa tatcggctgg acgaagtgcc ccaacttttt aatgttttaa 1560 tgggggatat gagctttgta ggtgtaagac cagaagtaca aaaatatgta aatcagtata 1620 ctgatgaaat gtttgcgacg ttacttttac ctgcaggaat tacttcacca gcgagtattg 1680 catataagga tgaagatatt gttttagaag aatattgttc tcaaggctat agtcctgatg 1740

aagcatatgt tcaaaaagta ttaccagaaa aaatgaagta caatttggaa tatatcagaa 1800 actttggaat tatttctgat tttaaagtaa tgattgatac agtaattaaa gtaataaaat 1860 aggagattaa aatgacaaaa agacaaaata ttccattttc accaccagat attacccaag 1920 ctgaaattga tgaagttatt gacacactaa aatctggttg gattacaaca ggaccaaaga 1980 caaaagagct agaacgtcgg ctatcagtat ttacaggaac caataaaact gtgtgtttaa 2040 attetgetae tgeaggattg gaactagtet taegaattet tggtgttgga eeeggagatg 2100 aagttattgt teetgetatg acetatactg ceteatgtag tgteattact eatgtaggag 2160 caactcctgt gatggttgat attcaaaaaa acagctttga gatggaatat gatgctttgg 2220 aaaaagcgat tactccgaaa acaaaagtta tcattcctgt tgatctagct ggtattcctt 2280 gtgattatga taagatttat accatcgtag aaaacaaacg ctctttgtat gttgcttctg 2340 ataataaatg gcagaaactt tttgggcgag ttattatcct atctgatagt gcacactcac 2400 taggtgctag ttataaggga aaaccagcgg gttccctagc agattttacc tcattttctt 2460 tccatgcagt taagaatttt acaactgctg aaggaggtag tgtgacatgg agatcacatc 2520 ctgatttgga tgacgaagag atgtataaag agtttcagat ttactctctt catggtcaga 2580 caaaggatgc attagctaag acacaattag ggtcatggga atatgacatt gttattcctg 2640

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gactagaaaa atgatgtaca attacggcta tccagggtgt ttgactttca tgtatgatgc 3600

agacaaaatg ggtttaattc agataaaaga tataaagaaa aataacgatt atgcgatatt 3660

acttcaattg tgtaagaagt atgactgtta tcttttaaat gaaagtttag cttcgtatcg 3720

aattagaaaa aaatcgat

3738

- <210> 44
- <211> 238
- <212> PRT
- <213> Streptococcus suis
- <220>
- <221> misc\_feature
- <223> CPS7E
- <400> 44

Ala Ala His Lys His Val Pro Leu Met Glu Tyr Asn Pro His Glu Ala

- 1
- 5
- 10
- 15

| Val Lys Asn | Asn Ile Phe Gly | Thr Lys Asn Va | ıl Ala Glu Ala Ala Lys |
|-------------|-----------------|----------------|------------------------|
| 20          | 25              | 30             |                        |

| Gly Glu Lys Leu | Tyr Glu Glu | Leu Leu Ser | Thr Glu | Glu Arg | Val Sei |
|-----------------|-------------|-------------|---------|---------|---------|
| 180             | 185         | 190         |         |         |         |

## $Asn\ Glu\ Leu\ Lys\ Asp\ Met\ Leu\ Ile\ Glu\ Phe\ Ala\ Lys\ Gln\ Glu$

<220>

| Met Thr Arg Val Glu Leu Ile Thr Arg Glu Phe Phe Lys Lys Asn Glu |                 |                       |                     |  |
|---|-----------------|-----------------------|---------------------|--|
| 1 5   | 10              | . 15                  |                     |  |
|   |                 |                       |                     |  |
| Ala Thr Ser I   | Lys Tyr Phe Gl  | n Lys Ile Glu Ser Ar  | g Arg Gly Glu Leu   |  |
| 20  | 25              | 30                    |                     |  |
|   |                 |                       |                     |  |
|   |                 | p Lys Leu Leu Ala I   | Leu Ile Leu Leu Leu |  |
| 35  | 40              | 45                    |                     |  |
|   | D 17 1 11 11    | TI T AL TI OD T       |                     |  |
|   |                 | Ile Leu Ala Ile Trp I | le Lys Leu Asp      |  |
| 50  | 55              | 60                    |                     |  |
|   |                 |                       |                     |  |
| Ser Lys Gly F   | Pro Ile Phe Tyr | Arg Gln Glu Arg V     | al Thr Arg Tyr Gly  |  |
| 65  | 70              | 75 80                 |                     |  |
|   |                 |                       |                     |  |
| Arg Ile Phe A   | arg Ile Phe Lys | Phe Arg Thr Met Ile   | e Ser Asp Ala Asp   |  |
| 85  | 90              | 95                    |                     |  |
|   |                 |                       |                     |  |
| Lys Val Gly S   | Ser Leu Val Th  | r Val Gly Gln Asp A   | sn Arg Ile Thr Lys  |  |
| 100   | 105             | 110                   |                     |  |
|   |                 |                       |                     |  |
| Val Gly His Ile Ile Arg Lys Tyr Arg Leu Asp Glu Val Pro Gln Leu |                 |                       |                     |  |
| 115   | 120             | 125                   |                     |  |
|   |                 |                       |                     |  |
| Phe Asn Val Leu Met Gly Asp Met Ser Phe Val Gly Val Arg Pro Glu |                 |                       |                     |  |
| 130   | 135             | 140                   |                     |  |
|   |                 |                       |                     |  |

Val Gln Lys Tyr Val Asn Gln Tyr Thr Asp Glu Met Phe Ala Thr Leu

Leu Leu Pro Ala Gly Ile Thr Ser Pro Ala Ser Ile Ala Tyr Lys Asp

165

170

175

Glu Asp Ile Val Leu Glu Glu Tyr Cys Ser Gln Gly Tyr Ser Pro Asp

180

185

190

Glu Ala Tyr Val Gl<br/>n Lys Val Leu Pro Glu Lys Met Lys Tyr As<br/>n Leu

195

200

205

Glu Tyr Ile Arg Asn Phe Gly Ile Ile Ser Asp Phe Lys Val Met Ile

210

215

220

Asp Thr Val Ile Lys Val Ile Lys

225

230

<210> 46

<211> 404

<212> PRT

<213> Streptococcus suis

<220>

<221> misc\_feature

<223> CPS7G

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1 5 10 15

Ala Glu Ile Asp Glu Val Ile Asp Thr Leu Lys Ser Gly Trp Ile Thr
20 25 30

Thr Gly Pro Lys Thr Lys Glu Leu Glu Arg Arg Leu Ser Val Phe Thr 35 40 45

Gly Thr Asn Lys Thr Val Cys Leu Asn Ser Ala Thr Ala Gly Leu Glu
50 55 60

Leu Val Leu Arg Ile Leu Gly Val Gly Pro Gly Asp Glu Val Ile Val
65 70 75 80

Pro Ala Met Thr Tyr Thr Ala Ser Cys Ser Val Ile Thr His Val Gly
85 90 95

Ala Thr Pro Val Met Val Asp Ile Gln Lys Asn Ser Phe Glu Met Glu
100 105 110

Tyr Asp Ala Leu Glu Lys Ala Ile Thr Pro Lys Thr Lys Val Ile Ile 115 120 125

Pro Val Asp Leu Ala Gly Ile Pro Cys Asp Tyr Asp Lys Ile Tyr Thr
130 135 140

Ile Val Glu Asn Lys Arg Ser Leu Tyr Val Ala Ser Asp Asn Lys Trp Gln Lys Leu Phe Gly Arg Val Ile Ile Leu Ser Asp Ser Ala His Ser Leu Gly Ala Ser Tyr Lys Gly Lys Pro Ala Gly Ser Leu Ala Asp Phe Thr Ser Phe Ser Phe His Ala Val Lys Asn Phe Thr Thr Ala Glu Gly Gly Ser Val Thr Trp Arg Ser His Pro Asp Leu Asp Asp Glu Glu Met Tyr Lys Glu Phe Gln Ile Tyr Ser Leu His Gly Gln Thr Lys Asp Ala Leu Ala Lys Thr Gln Leu Gly Ser Trp Glu Tyr Asp Ile Val Ile Pro Gly Tyr Lys Cys Asn Met Thr Asp Ile Met Ala Gly Ile Gly Leu Val 

Gln Leu Glu Arg Tyr Pro Ser Leu Leu Asn Arg Arg Glu Ile Ile 275 280 285

Glu Lys Tyr Asn Ala Gly Phe Glu Gly Thr Ser Ile Lys Pro Leu Val 290 295 300

| His Leu Thr Glu Asp Lys Gln Ser Ser Met His Leu Tyr Ile Thr His |                |              |                  |                    |  |
|---|----------------|--------------|------------------|--------------------|--|
| 305   | 310            | 315          | 320              |                    |  |
|   |                |              |                  |                    |  |
| Leu Gln   | Gly Tyr Thr l  | Leu Glu Gln  | Arg Asn Glu Va   | al Ile Gln Lys Met |  |
|   | 325            | 330          | 335              |                    |  |
|   |                |              |                  |                    |  |
| Ala Glu   | Ala Gly Ile Al | la Cys Asn V | al His Tyr Lys l | Pro Leu Pro Leu    |  |
| 3   | 340            | 345          | 350              |                    |  |
|   |                |              |                  |                    |  |
| Leu Thr Ala Tyr Lys Asn Leu Gly Phe Glu Met Lys Asp Phe Pro Asn |                |              |                  |                    |  |
| 355   | 36             | 50           | 365              |                    |  |
|   |                |              |                  |                    |  |
| Ala Tyr   | Gln Tvr Phe (  | Hu Asn Glu   | Val Thr Leu Pro  | Leu His Thr Asn    |  |

Ala Tyr Gln Tyr Phe Glu Asn Glu Val Thr Leu Pro Leu His Thr Asn 370 375 380

Leu Ser Asp Glu Asp Val Glu Tyr Val Ile Glu Met Phe Leu Lys Ile 385 390 395 400

Val Ser Arg Asp

<210> 47

<211> 210

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<213> Streptococcus suis

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| 77     | 7  | "  | ` |
| $\sim$ | Z. | ١, | _ |

<221> misc\_feature

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20 25 30

Val Leu Asp Gln Thr His Gln Asn Trp Glu Leu Ile Ile Val Asp Asp 35 40 45

Cys Ser Asn Asp Glu Thr Glu Lys Val Val Ser His Phe Lys Asp Ser 50 55 60

Arg Ile Lys Phe Phe Lys Asn Ser Asn Asn Leu Gly Ala Ala Leu Thr
65 70 75 80

Arg Asn Lys Ala Leu Arg Lys Ala Arg Gly Arg Trp Ile Ala Phe Leu 85 90 95

Asp Ser Asp Asp Leu Trp His Pro Ser Lys Leu Glu Lys Gln Leu Glu
100 105 110

| Phe Met Lys A | sn Asn Gly | Гуг Ser Phe Thr | Tyr His Asn Phe Glu Lys |
|---------------|------------|-----------------|-------------------------|
| 115           | 120        | 125             |                         |

Lys Lys

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<211> 101

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- <223> N may be any nucleotide
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- caancatttt aaattttaga aaattagttt ttagagctcc c
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- <220>
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- <400> 49
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- caancatett aaattttaga aaattagttt ttagaggtee e
- 101

- <210> 50
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- <220>
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# <223> 100 base pair repeat between CPS2O and CPS2P

<400> 50

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caaacatttt aaattttaga aaattagttt ttagaggtcc c

101

<210> 51

<211> 120

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<213> Streptococcus suis

<220>

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<400> 51

 $\label{eq:metalaction} \mbox{Met Ala Lys Val Ser Ile Ile Val Pro Ile Phe Asn Thr Glu Lys Tyr}$ 

1 5

10

15

Leu Arg Glu Cys Leu Asp Ser Ile Ile Ser Gln Ser Tyr Thr Asn Leu

20

25

30

| Glu Ile Leu Leu Ile Asp Asp Gly | Ser Ser Asp Ser Ser Thr Asp 1 | [le |
|---------------------------------|-------------------------------|-----|
|---------------------------------|-------------------------------|-----|

35

40

45

Cys Leu Glu Tyr Ala Glu Gln Asp Gly Arg Ile Lys Leu Phe Arg Leu 50 55 60

Pro Asn Gly Gly Val Ser Asn Ala Arg Asn Tyr Gly Ile Lys Asn Ser
65 70 75 80

Thr Ala Asn Tyr Ile Met Phe Val Asp Ser Asp Asp Ile Val Asp Gly
85 90 95

Asn Ile Val Glu Ser Leu Tyr Thr Cys Leu Lys Glu Asn Asp Ser Asp 100 105 110

Leu Ser Gly Gly Leu Leu Ala Thr

115

120

<210> 52

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- <400> 52

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Leu Ser Lys Cys Ile Asn Ser Ile Val Asn Gln Thr Tyr Lys His Ile

Glu Leu Leu Val Asn Asp Gly Ser Ser Thr Asp Asn Ser Glu Glu Ile

Cys Leu Ala Tyr Ala Lys Lys Asp Ser Arg Ile Arg Tyr Phe Lys Lys

Glu Asn Gly Gly Leu Ser Asp Ala Arg Asn Tyr Gly Ile Ser Arg Ala

Lys Gly Asp Tyr Leu Ala Phe Ile Asp Ser Asp Asp Phe Ile His Ser

- 100
- 105
- 110

## Leu Xaa Xaa Val Ala Val Ala Gly

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- 120
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- <220>
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- <400> 53

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- 1
- 5
- 10
- 15

Glu His Arg Phe Lys Arg Gly Glu Lys Leu Pro Ser Ile Arg Gln Leu

- 20
- 25
- 30

| Arg Glu Gln Tyr His Cys Ser Lys Asp Thr Val Gln Lys Ala Met Leu |                 |             |                       |           |
|---|-----------------|-------------|-----------------------|-----------|
| 35  | 40              | 45          |                       |           |
|   |                 |             |                       |           |
| Glu Leu Lys   | Гуг Gln Asn Ly  | s Ile Tyr A | Ala Val Glu Lys Ser   | Gly Tyr   |
| 50  | 55              | 60          |                       |           |
|   |                 |             |                       |           |
| Tyr Ile Leu G   | lu Asp Arg As   | p Phe Gln   | Asp His Thr Cys Ar    | g Ala Gln |
| 65  | 70              | 75          | 80                    |           |
|   |                 |             |                       |           |
| Ser Tyr Arg I   | Leu Ser Arg Ile | Thr Tyr C   | Glu Asp Phe Arg Ile   | Cys Leu   |
| 85  | 90              |             | 95                    |           |
|   |                 |             |                       |           |
| Lys Glu Ser L   | eu Ile Gly Arg  | Glu Asn T   | Гуг Leu Phe Asn Туг   | Tyr His   |
| 100   | 105             | 1           | 10                    |           |
|   |                 |             |                       |           |
| Gln Gln Glu C   | Gly Leu Ala Gl  | u Leu Ile S | Ser Ser Val Gln Ser I | Leu Leu   |
| 115   | 120             | 125         |                       |           |
|   |                 |             |                       |           |
| Met Asp Tyr His Val Tyr Thr Lys Lys Asp Gln Leu Val Ile Thr Ala |                 |             |                       |           |
| 130   | 135             | 140         |                       |           |
|   |                 |             |                       |           |
| Gly Ser Gln Gln Ala Leu Tyr Ile Leu Thr Gln Met Glu Thr Leu Ala |                 |             |                       |           |
| 145   | 150             | 155         | 160                   |           |
|   |                 |             |                       |           |

Glu Leu Ile Arg His Gln Gly Ile Pro Tyr Gln Thr Ile Glu Arg Asn 180 185 190

Gly Lys Thr Glu Ile Leu Ile Glu Asn Pro Thr Tyr Ser Arg Met Ile

175

170

165

Leu Asp Gly Ile Asp Leu Glu Glu Leu Glu Ser Ile Phe Gln Thr Gly 

Lys Ile Lys Phe Phe Tyr Thr Ile Pro Arg Leu His Asn Pro Leu Gly 

Ser Thr Tyr Asp Ile Ala Thr Lys Thr Ala Ile Val Lys Leu Ala Lys 

Gln Tyr Asp Val Tyr Ile Ile Glu Asp Asp Tyr Leu Ala Asp Phe Asp 

Ser Ser His Ser Leu Pro Leu His Tyr Leu Asp Thr Asp Asn Arg Val 

Ile Tyr Ile Lys Ser Phe Thr Pro Thr Leu Phe Pro Ala Leu Arg Ile 

Gly Ala Ile Ser Leu Pro Asn Gln Leu Arg Asp Ile Phe Ile Lys His 

Lys Ser Leu Ile Asp Tyr Asp Thr Asn Leu Ile Met Gln Lys Ala Leu 

Ser Leu Tyr Ile Asp Asn Gly Met Phe Ala Arg Asn Thr Gln His Leu 

His His Ile Tyr His Ala Gln Trp Asn Lys Ile Lys Asp Cys Leu Glu 

Lys Tyr Ala Leu Asn Ile Pro Tyr Arg Ile Pro Lys Gly Ser Val Thr 

Phe Gln Leu Ser Lys Gly Ile Leu Ser Pro Ser Ile Gln His Met Phe 

Gly Lys Cys Tyr Tyr Phe Ser Gly Gln Lys Ala Asp Phe Leu Gln Ile 

Phe Phe Glu Gln Asp Phe Ala Asp Lys Leu Glu Gln Phe Val Arg Tyr 

Leu Asn Glu

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#### **CLAIMS**

#### What is claimed is:

- 1. An isolated or recombinant nucleic acid encoding a capsular gene cluster of Streptococcus suis or a gene or gene fragment derived thereof.
- 2. The isolated or recombinant nucleic acid of claim 1, wherein said nucleic acid encodes a *Streptococcus suis* serotype-specific central region.
- 3. The isolated or recombinant nucleic acid of claim 1 or claim 2, wherein said isolated or recombinant nucleic acid is hybridized to a second nucleic acid encoding a gene derived from a *Streptococcus suis* serotype 1, 2, or 9 capsular gene cluster.
- An isolated or recombinant nucleic acid encoding a capsular gene cluster of *Streptococcus suis* serotype 2 or a gene or gene fragment derived thereof, wherein said isolated or recombinant nucleic acid comprises SEQ. ID. NO. 9 and said isolated or recombinant nucleic acid encodes a capsular gene cluster of *Streptococcus suis* serotype 2 or a gene or gene fragment derived thereof selected from the group of sequences consisting of SEQ. ID. NO. 10, SEQ. ID. NO. 53, SEQ. ID. NO.11, SEQ. ID. NO.12, SEQ. ID. NO.13, SEQ. ID. NO.14, SEQ. ID. NO.15, SEQ. ID. NO.16, SEQ. ID. NO.17, SEQ. ID. NO.18, SEQ. ID. NO.19, SEQ. ID. NO.20, SEQ. ID. NO.21, SEQ. ID. NO.22, SEQ. ID. NO.23, SEQ. ID. NO.24, SEQ. ID. NO.25, SEQ. ID. NO.26, SEQ. ID. NO.27 and SEQ. ID. NO. 28.
- 5. An isolated or recombinant nucleic acid encoding a capsular gene cluster of *Streptococcus suis* serotype 1 or a gene or gene fragment derived thereof, wherein said isolated or recombinant nucleic acid is SEQ. ID. NO.29 and said isolated or recombinant nucleic acid encodes a capsular gene cluster of *Streptococcus suis* serotype 1 or a gene or gene fragment derived thereof selected from the group consisting of SEQ. ID. NO.30, SEQ. ID. NO. 31, SEQ. ID. NO.32, SEQ. ID. NO.33, SEQ. ID. NO.34, SEQ. ID. NO.35 and SEQ. ID. NO.36.

- 6. An isolated or recombinant nucleic acid encoding a capsular gene cluster of *Streptococcus suis* serotype 9 or a gene or gene fragment derived thereof, wherein said nucleic acid comprises SEQ. ID. NO.37 and wherein said isolated or recombinant nucleic acid encodes a capsular gene cluster of *Streptococcus suis* serotype 9 or a gene or gene fragment derived thereof selected from the group consisting of SEQ. ID. NO.38, SEQ. ID. NO.39, SEQ. ID. NO.40, SEQ. ID. NO.41, and SEQ. ID. NO.42.
- 7. A nucleic acid probe or primer derived from the isolated or recombinant nucleic acid of any one of claims 1 to 6, wherein said nucleic acid probe or primer allows species or serotype specific detection of *Streptococcus suis*.
- 8. The nucleic acid probe or primer of claim 7, wherein said nucleic acid probe or primer further comprises at least one reporter molecule.
- 9. A diagnostic test kit comprising the nucleic acid probe or primer of claim 7 or claim 8.
- 10. A protein or fragment thereof encoded by the isolated or recombinant nucleic acid of any one of claims 1 to 6.
- 11. The protein or fragment of claim 10, wherein said protein or fragment is capable of polysaccharide biosynthesis.
- 12. A process for producing a *Streptococcus suis* capsular antigen, said method comprising:

using the protein or fragment of claim 11 to prepare said Streptococcus suis capsular antigen.

13. A Streptococcus suis capsular antigen produced by the process of claim 12.

### 14. A vaccine comprising:

the *Streptococcus suis* capsular antigen of claim 13 in an amount sufficient to produce an immune response in a subject, and

a suitable carrier or adjuvant.

- 15. A recombinant *Streptococcus suis* mutant having a modified capsular gene cluster.
- 16. A recombinant microorganism comprising at least a part of a capsular gene cluster of *Streptococcus suis*, wherein said <u>capsular</u> gene cluster comprises a deletion, insertion or (point)-mutation.
- 17. The recombinant microorganism of claim 16, wherein said recombinant microorganism comprises a lactic acid bacterium.
- 18. A vaccine comprising the recombinant *Streptococcus suis* mutant of claim 15 or the microorganism of claim 16 or claim 17.
- 19. The vaccine of claim 18, wherein said vaccine [comprises a Streptococcus mutant] includes a Streptococcus mutant deficient in capsular expression.
- 20. The vaccine of claim 19, wherein said Streptococcus mutant deficient in capsular expression is a recombinant Streptococcus mutant.
- 21. The vaccine of claim 19 or claim 20, wherein said Streptococcus mutant deficient in capsular expression is capable of surviving in an immune-competent host.
- 22. The vaccine of claim 21, wherein said Streptococcus mutant deficient in capsular expression is capable of surviving at least 4-5 days in said immune-competent host.

- 23. The vaccine of any one of claims 19 to 22, wherein said Streptococcus mutant deficient in capsular expression expresses a *Streptococcus* virulence factor or antigenic determinant.
- 24. The vaccine of any of claims 19 to 23, wherein said Streptococcus mutant deficient in capsular expression expresses a *non-Streptococcus* protein.
- 25. The vaccine of claim 24 wherein said non- *Streptococcus* protein has been derived from a pathogen.
- 26. A method for controlling or eradicating a Streptococcal disease in a population, said method comprising:

vaccinating subjects in said population with the vaccine of any one of claims 18 to 25.

A method for controlling or eradicating a Streptococcal disease, said method comprising:

testing for the presence of encapsulated Streptococcal strains in a sample collected from at least one subject in a population partly or wholly vaccinated with a vaccine of any one of claims 18 to 25.

A method for controlling or eradicating a Streptococcal disease comprising testing for the presence of capsule-specific antibodies directed against Streptococcal strains in a sample collected from at least one subject in a population partly or wholly vaccinated with a vaccine of any one of claims 18 to 25.

- 29. A method for controlling or eradicating a Streptococcal disease in a population comprising:
  - selecting subjects in said population vaccinated with a vaccine according to any one of claims 18 to 25; and
  - testing a sample collected from at least one subject in said population for the presence of encapsulated Streptococcal strains and/or for the presence of capsule-specific antibodies directed against Streptococcal strains.

### ABSTRACT OF THE DISCLOSURE

The invention relates to *Streptococcus suis* infection in pigs, vaccines directed against those infections and tests for diagnosing *Streptococcus suis* infections. The invention provides an isolated or recombinant nucleic acid encoding a capsular gene cluster of *Streptococcus suis* or a gene or gene fragment derived thereof. The invention further provides a nucleic acid probe or primer allowing species or serotype-specific detection of *Streptococcus suis*. The invention also provides a *Streptococcus suis* antigen and vaccine derived thereof.

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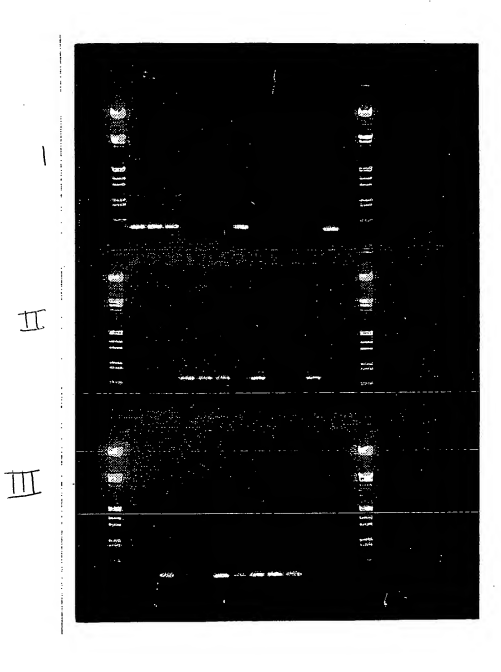


Fig.

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| ллссттссат  | ΔΤΤΓΩΤΓΆΓΑ | TGATGGAGGT | GATGGAAGCA               | TCTAAGTCTG          | CAGCGGGGTC           |
| CCCCTCCCCA  | AGTCCGCAGG | CTTATCAGGC | AGCTTTTGAG               | GGAGCTGAGA          |                      |
| ACATTATCGT  |            | ACAGGTGGGC | TATCGGGTAG               | TTTTAATGCG          | GCACGTGTAG           |
| CTAGGGATAT  |            |            | ATGTCAATAT               |                     |                      |
|   | CAGCCAGTGG |            |                          | ACCAAATCAA          | TCGCTTAATT           |
|   |            | ACABGTAGTA | GAAGCGATAA               |                     |                      |
| AGTGCAGGAT  | AAGCTCCTCT |            |                          | AATCTTGTTA          | AGAATGGAAG           |
|   | TTGGTAGGCA |            |                          | ATCCGTATGG          |                      |
|   | AAGTGCTGAA |            |                          | AAAGGCGCGT          | GGTCATAAGA           |
| TIGGIGAGGC  | ACCACCCTTT | GAAGAAATGA | AAAAAGCAGG               |                     |                      |
|   | TTATGGCCCA |            |                          | TCCAACAATT          | CTCAGAGTTG           |
|   | GTTTTCCAAC |            |                          | CAACATCAGG          |                      |
| TCTATGCAGT  |            | AAGAAGGTGG | ACTTTTGATG               |                     | TGAAAGCGTG           |
| ATTCACAGAG  |            | GGGCTGTAAT |                          | GAATAATCCC          |                      |
|   | TAAGTTCGAG |            |                          | TATTGGATTT          | CATTCATTCA           |
| מתידים של מידע מידע מידע מידע מידע מידע מידע מידע | GAATTGCTCC | AGTTTATCTG |                          | TTCAAAGAAG          |                      |
| AMIMICITADED                                      | AATCAGCTTT | CTGTCCGCTG | AAATAATAAC               | ATTTTCCAAA          | CATGTGTTGG           |
|   | AAAGAATCCC |            | TGAAAGGTCA               | CGCTCCCCTT          |                      |
|   | TACGGGATGT |            |                          | CAGTCTTTTA          | TTTTATTCCA           |
|   | TAAATGTGAT |            |                          | GCAAACATAC          |                      |
|   | GTAGAGCGAG |            |                          | ATTGGTATCG          | TAGTCGATTA           |
| GACTCTTATG  | TTTGATGAAG | ATATCACGTA | GCTGATTAGG               | AAGGCTGATT          |                      |
| GCACCGATTC  | GGAGGGCAGG |            | GGTGTAAAAG               | ATTTTATATA          | GATGACGCGA           |
| TTATCTGTAT  | CAAGATAGTG | TAAAGGTAGG | CTATGACTAG               |                     |                      |
| TGCTAAATAG  | TCATCCTCAA | TGATGTAGAC | ATCGTATTGC               | TTTGCTAATT          | TTACGATGGC           |
|   | GCTATATCAT |            | GAGAGGGTTG               | TGCAAGCGAG          |                      |
| GAATTGTGTA  | GAAAAACTTA | ATTTTTCCAG | TTTGGAAGAT               | ACTTTCCAAT          | TCTTCTAGGT           |
| CAATTCCATC  | TAAATTCCGT | TCAATTGTTT |                          | TCCTTGATGT          |                      |
| CGAATGAGCT  | CTATCATTCG | TGAATAGGTA | GGGTTCTCTA               | TCAAGATTTC          | CGTTTTTCCA           |
| GCCAAGGTTT  | CCATTTGTGT | GAGAATATAT | AGAGCTTGTT               | GACTACCAGC          |                      |
| TGTGATAACC  | AGCTGGTCTT | TTTTTGTATA | GACATGATAG               |                     | GACTTTGAAC           |
|   | AATTCTGCCA |            | CTGGTGATAG               | TAGTTGAATA          |                      |
|   | CCGCCCAATA |            | TTAGACAAAT               | CCGAAAATCT          | TCATAGGTAA           |
|   | TCTGTAGGAT |            | AGGTATGGTC               | TTGGAAATCT          |                      |
|   | AGATATAATA |            |                          | AGATCTTATT          | TTGGTATTTT           |
|   | TAGCCTTTTG |            | TTGCTACAAT               | GATATTGCTC          |                      |
|   | CGGATAGAAG |            |                          | AATCGATGTT          | CCTCTATTCC           |
|   | TCTTGGATGA |            | TTTTTTCATC               |                     | CARMAMAMCA           |
| TTTTTATAGA  | CTATGTTACT | AGCTAGTATA |                          | TGAAGAAAGA          | CAATATATGA           |
| ATAATGGGGT  | TGAGGTTCAG | GAATTAAGCT | ACTCTATGGT               | ATAATTAAGT          | A M.C.C.C.M.C.C.M.M. |
| GATGAAAATA  | ATTATACCTA | ATGCAAAAGA | AGTAAATACA               | AATCIAGAGA          | AIGCCICGII           |
| TTATCTCCTG  | TCTGATCGAA | GCAAGCCGGT | GCTGGATGCC               | AIAMGICAMI          | САСТТАСААС           |
| TTGATGTAAA  | AAAGATGGCT | GCCTTTTATA | AATTGAATGA               | CTATCCACCC          | GAGIIAGAAG           |
| CTGACCGTTG  | GTATCGAATC | AGGACAGGTC | AAGCAAAAAC               | CCCCACCTAT          | AGATTCGAAA           |
| TGGCAGTTAT  | ATGATGGTCT | CATGTATCGT | CERCCCACAC               | CCTTATACCC          | AGATTCGAAA           |
| GAAGAAAATT  | ATTTACGTGA | CCACGITCGI | GTAGCGACAG               | CATTTTCAAC          | CCACCTTAAA           |
| ATTGATTCAT  | CCTTTTGAAT | TCATTTCACC | CCCACCCTAT               | TATCACCAAG          | GGAGCTTAAA           |
| GATAGGCAAT  | CAGTCTTTGA | AACAGTACTG | GCGACCGTAT<br>TGGCTTCGTC | ACAAMMTCAC          | СХССТСТТТТ           |
| AAGTTGGTGA  | TGATGAACTG | ATTCTCTCAC | TTCTTTTCAT               | CCANCADAA           | CAGGIGITI            |
| CTCCCCAGAT  | TCAGAAAAGA | TTAGTTAAAA | TICITIONI                | CCACACCAAC          | <b>ΔΨΨ</b> ΘΟΨΩΨΟΟ   |
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| AATTGGTAAA  | TTTTGCGCTT | TTGGGACTTT | ATTCCATTAC               | TOTATOTITO          | IICIIAGIGA           |
| CCATGTATCG  | CTATAACATC | CTAGATTTCC | GGTATTTAAA               | TATALIGIO COCO      | TARGARGE             |
| ACGCTTTTGC  | TAGTAGGAGT | GGCAGTATTG | GCTGGATTAT               | TOWIGIOGCO          | . ANGANAGCO          |
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TGGGATCTAT GGAATGCAAG AAGTTGTAAA ATTTTCAACA CGACTAAATT CAAATTCGAC ATTTTCAGAA TATGAAATGA GTATCCTTGT CCCAGCAAAT AGTGATATTA CGGACGTTCG TCAGCTTACT AGTATCCTTG CTCCAGCCGA ATACGACCAA GATAACATCA CCGCTTTATT GGATGACATA TCCAAAATGG AATCTACTCA ACTAGCAACT AGCCCCGGGA CTTCTTACCT GACAGCATAT CAATCTATGT TGAATGGCGA GAGTCAAGCG ATGGTGTTCA ACGGAGTTTT TACCAATATT TTAGAAAATG AAGATCCAGG CTTTTCTTCA AAAGTGAAAA AAATATATAG TTTCAAAGTG ACTCAGACTG TTGAAACAGC TACTAAGCAG GTGAGTGGAG ATAGCTTTAA TATCTATATT AGTGGTATTG ATGCTTATGG ACCGATTTCT ACGGTCTCTC GTTCAGATGT CAATATCATT ATGACTGTCA ATCGTGCGAC ACATAAGATT TTATTGACAA CTACTCCACG AGATTCATAC GTTGCTTTCG CAGATGGCGG GCAAAATCAA TACGATAAAC TAACACATGC TGGTATTTAC GGTGTCAATG CTTCTGTGCA CACCTTAGAA AATTTTTATG GGATTGACAT TAGGATIAI GTGCGGTTGA ACTTCATTTC CTTCCTTCAA TTAATCGACT TGGTGGGTGG

AATTGATGTA TATAACGATC AAGAATTTAC AAGTTTACAT GGGAATTATC
ATTTCCCTGT TGGACAAGTT CATTTAAACT CAGACCAAGC ATTAGGCTTC GCTACTCTTT AACAGGGGGT GACAATGACC GTGGTAAAAA CCAGGAAAAA TTACCAGGCA ATCTAAAAAA TTACCAGGCA ATCTAAAAAA TTACCAGGCA ATCTAAAAAA TTACCAGGCA ATCTAAAAAA TTACCAGGCA ATCTAAAAAA TTACCAGGCA ACCATTGACA ACGATTTGA GCTTAGAAAC CAATTTACAG GATCACAACT TTATATGATG GAAATTAACC AAGATAGTCT GCGATGCCTG GAGCAATCA AAGATAAACA TTATAGAGA ACTTGTTGAA AAATAAAGAT TTTAGGAGAA AATATGAACA AATTCCAGTA AAATCCAATC GAAATCCATC GAAATCCATC GAAATCCATC GAAATCCATC TTATATGACA AAATCCAATC TTATATCTT ACTGAGAAA AATTCCATT TTATATCTT ACTGAGAAA AATTCCATT TTATATCTT TTATGGAGAA AATTTCCATTA ACTGCAGTGT TGACTGCGGG GTTGGCATTT TTATGGAGAA ATTTGGAGAA ATTTGAGAA ATTTGGAGAA ATTTGAA ATTTGGAGA TAGCAATTAT GTGCGGTTGA ACTTCATTTC CTTCCTTCAA TTAATCGACT TGGTGGGTGG GTCTACAGTA GTTTTTAGT GACACCTCAA TATGACTCCA CTACCCGTAT CTATGTAGTG AGTCAAAATG TTGAAGCCGG TGCGGGCTTG ACTAACCAAG AGTTACAAGC GGGTACCTAT TTGGCAAAAG ACTATCGGGA AATTATCCTA TCACAAGATG TATTGACACA AGTAGCAACG GAATTGAATC TGAAAGAGA TTTGAAAGAA AAAATATCAG TTTCTATTCC TGTTGATACT CGTATCGTTT CTATTTCTGT GCGTGATGCG GATCCAAATG → AAGCGGCACG TATTGCAAAT AGCCTTCGCA CCTTTGCAGT GCAAAAGGTT GTTGAGGTCA CCAAGGTAAG CGATGTGACG ACACTTGAAG AAGCAGTCCC AGCGGAAGAA CCAACCACTC CAAATACAAA ACGAAATATC TTGCTTGGTT TATTAGCTGG AGGTATCTTG GCAACAGGTC TTGTACTGGT TATGGAGGTT TTGGATGACC GTGTAAAACG TCCTCAGGAC ATCGAAGAGG TAATGGGATT GACATTGCTA GGTATAGTAC CAGATTCGAA GAAATTAAAA TAGGAGAACA ATATGGCGAT GTTAGAAATT GCACGTACAA AAAGAGAGGG AGTAAATAAA ACCGAGGAGT ATTTCAATGC TATCCGTACC AATATTCAGC TTAGCGGAGC AGATATTAAG GTTGTTGGTA TTACCTCTGT TAAATCGAAT GAAGGTAAGA GTACAACTGC GGCTAGTCTC GCTATTGCCT ATGCTCGTTC AGGTTATAAG ACCGTCTTGG TGGATGCAGA TATCCGAAAT TCAGTCATGC CTGGTTTCTT CAAGCCAATT ACAAAGATTA CAGGTTTGAC GGATTACCTA GCAGGGACAA CAGACTTGTC TCAAGGATTA TGCGATACAG ATATTCCAAA CTTGACCGTA ATTGAGTCAG GAAAGGTTTC TCCCAACCCT ACTGCCCTTT TACAAAGTAA GAATTTTGAA AATCTACTTG CGACTCTTCG TCGCTATTAT GATTATGTTA TCGTTGACTG TCCACCATTA GGACTGGTAA TTGATGCAGC TATCATTGCA CAAAAATGTG ATGCGATGGT TGCAGTAGTA GAAGCAGGCA ATGTTAAGTG CTCATCTTTG AAAAAAGTAA AAGAGCAGTT GGAACAAACA GGCACACCGT TCTTAGGCGT TATCTTGAAC AAATATGATA TTGCCACTGA GAAGTATAGT GAATACGGAA ATTACGGCAA AAAAGCCTAA TTTCTCAGAT AACATAAGTT TGATAAGTAG GTATTAATAT GATTGATATC CATTCGCATA TCATATTTGG TGTGGATGAC GGTCCCAAAA CTATTGAAGA GAGCCTGAGT TTGATAAGCG AAGCTTATCG TCAAGGTGTT CGCTATATCG TAGCGACATC TCATAGACGA AAAGGGATGT TTGAAACACC AGAAAAAATC ATCATGATTA ACTTTCTTCA ACTTAAAGAG GCAGTAGCAG AAGTTTATCC TGAAATACGA TTGTGCTATG GTGCTGAATT GTATTATAGT AAAGATATCT TAAGCAAACT TGAAAAAAAG AAAGTACCAA CACTTAATGG CTCGTGCTAT ATTCTCTTGG AGTTCAGTAC GGATACTCCT TGGAAAGAGA TTCAAGAAGC AGTGAACGAA ATGACGCTAC TTGGGCTAAC TCCCGTACTT GCCCATATAG AGCGTTATGA TGCTCTGGCA TTTCAGTCAG AGAGAGTAGA AAAGCTAATT GACAAGGGAT GCTACACTCA GGTAAATAGT AACCATGTGT TGAAGCCTGC TTTAATTGGC GAACGAGCAA AAGAATTTAA AAAACGTACT CGATATTTTT TAGAGCAGGA TTTAGTACAT TGTGTTGCTA GCGATATGCA TAATTTATAT AGTAGACCTC CGTTTATGAG GGAGGCGTAT CAGCTTGTAA AAAAAGAGTA TGGTGAGGAT AGAGCGAAGG

Fig. 3 cont.

CTTTGTTCAA GAAAAATCCT TTGTTGATAT TGAAAAATCA AGTACAGTAA CCTCATAGAA ATAGTGGAGG AGCTATGAAT ATTGAAATAG GATATCGCCA AACGAAATTG GCATTGTTTG ATATGATAGC AGTTACGATT TCTGCAATCT TAACAAGTCA TATACCAAAT GCTGATTTAA ATCGTTCTGG AATTTTTATC ATAATGATGG TTCATTATTT TGCATTTTTT ATATCTCGTA TGCCGGTTGA ATTTGAGTAT AGAGGTAATC TGATAGAGTT TGAAAAACA TTTAACTATA GTATAATATT TGTAATTTTT CTTATGGCAG TTTCATTTAT GTTAGAGAAT AATTTCGCAC TTTCAAGACG TGGTGCCGTG TATTTCACAT TAATAAACTT CGTTTTGGTA TACCTATTTA ACGTAATTAT TAAGCAGTTT AAGGATAGCT TTCTATTTTC GACAACCTAT CAAAAAAAGA CGATTCTAAT TACAACGGCT AGTCGGTACC CGTCCGCCTA CAGTTGATGA ATTTGAAAAA TATACTCCTA GTCAAAAGAG AAGATTGAGT TTTAAACCAG GGATTACAGG TCTTTGGCAA GTGAGCGGAA GAAGTGATAT CACAGATTTT AATGAAGTCG TTAGGCTGGA CCTAACATAC ATTGATAATT GGACCATCTG GTCAGACATT AAGATTTTAT TGAAGACAGT GAAAGTTGTA TTGTTGAGAG AGGGAGGTCA GTAAGACTCC TTTAAAACAA AGAATAGTAG TAGGGGATAT GAGAACAGTT TATATTATTG GTTCAAAAGG AATACCAGCA AAGTATGGTG GTTTCGAGAC TTTCGTAGAA AAATTAACTG AGTATCAGAA AGATAAATCA ATTAATTATT TTGTTGCATG TACAAGAGAA AATTCAGCAA AATCAGATAT TACAGGAGAA GTTTTTGAAC ATAATGGAGC AACATGTTTT AATATTGATG TGCCAAATAT TGGTTCAGCA AAAGCCATTC TTTATGATAT TATGGCTCTC AAGAAATCTA TTGAAATTGC CAAAGATAGA AATGATACCT CTCCAATTTT CTACATTCTT GCTTGTCGGA TTGGTCCTTT CATTTATCTT TTTAAGAAGC AGATTGAATC AATTGGAGGT CAACTTTTCG TAAACCCAGA CGGTCATGAA TGGCTACGTG AAAAGTGGAG TTATCCCGTC CGACAGTATT GGAAATTTTC TGAGAGTTTG ATGTTAAAAT ACGCTGATTT ACTAATTTGT GATAGCAAAA ATATTGAAAA ATATATTCAT GAAGATTATC GAAAATATGC TCCTGAAACA TCTTATATTG CTTATGGAAC AGACTTAGAT AAATCACGCC TTTCTCCGAC AGATAGTGTA GTACGTGAGT GGTATAAGGA GAAGGAAATT TCAGAAAATG ATTACTATTT GGTTGTTGGA CGATTTGTGC CTGAAAATAA CTATGAAGTA ATGATTCGAG AGTTTATGAA ATCATATTCA AGAAAAGATT TTGTTTTGAT AACGAATGTA GAGCATAATT CCTTTTATGA GAAATTGAAA AAAGAAACAG GGTTCGATAA AGATAAGCGT ATAAAGTTTG TTGGAACAGT CTATAATCAG GAGCTGTTAA AATATATTCG TGAAAATGCA TTTGCTTATT TTCATGGTCA CGAGGTTGGA GGAACGAACC CATCTTTACT TGAAGCACTT TCTTCTACTA AACTAAATCT TCTTCTAGAT GTGGGCTTTA ATAGAGAAGT AGGGGAAGAA GGAGCGAAAT ACTGGAATAA AGATAATCTT CACAGAGTTA TTGACAGTTG TGAGCAATTA TCACAAGAAC AAATTAATGA TATGGATAGT TTATCAACAA AACAAGTCAA AGAAAGATTT TCTTGGGATT TTATTGTTGA TGAGTATGAG AAGTTGTTTA AAGGATAAGT TATGAAAAAG ATTCTATATC TCCATGCTGG AGCAGAATTA TATGGGGCAG ATAAGGTTCT CTTGGAACTT ATAAAAGGCT TAGATAAGAA TGAATTTGAA GCGCATGTTA TCCTACCTAA TGATGGAGTC CTAGTGCCAG CATTAAGAGA AGTTGGTGCG CAAGTTGAAG TTATTAACTA TCCAATTCTA CGTAGGAAAT TTGCTCAATA TGCCATAGAA AATAAGGTTG ACATAATTCA CAATAATACT ACCGCTGTCT TAGAAGGCAT TTATCTGAAG CGAAAACTCA AATTACCTTT GTTGTGGCAT GTTCATGAGA TTATTGTCAA ACCTAAATTC ATCTCTGATT CGATCAATTT TTTAATGGGG CGTTTTGCTG ATAAGATTGT GACAGTTTCA CAGGCTGTGG CAAACCATAT AAAACAATCA CCTCATATCA AAGATGACCA AATCAGTGTA ATCTACAATG GGGTAGATAA TAAAGTGTTT TATCAGTCCG ATGCTCGGTC TGTTCGAGAA AGATTTGACA TTGACGAAGA GGCTCTTGTC ATTGGTATGG TCGGTCGAGT CAATGCGTGG

## Charge Margens

6/59

| •                        | 1                        | 6/5                          | 59                                     | •           |                   |
|--------------------------|--------------------------|------------------------------|--|-------------|-------------------|
| AAAGGACAAG               | GAGATTTTTT               | AGAAGCAGTT                   | GCTCCTATAC                             | TCGAACAGAA  | TCCAAAAGCT        |
| ATCGCCTTTA               |                          | TGCTTTTGAA                   | GGAGAAGAGT                             | GGCGAGTAGT  |                   |
|                          | AAGAAGATTT               | CTCAATTAAA                   | GGTCTCTTCT                             | CAAGTCAGAC  | GAATGGATTA        |
|                          | ACCACTGAAT               |                              | GTTTGATATT                             |             |                   |
|                          | TCCAGACCCT               | CTACCAACGG                   | ТТСТАСТАВА                             | AGCAATGGCA  | TGCGGTAAAC        |
|                          |                          |                              | GTGAGATGGT                             |             |                   |
| CTGTTGTCGG               |                          |                              |  | TATCAAAAGT  | AATTCTTCAG        |
| GTTAACGGTT               | TCTTAGTCAC<br>ATATAAATCT |                              |  |             |                   |
| TTATCGGAAA               | GAACATTTTT               | CAGAAAAAAA                   | CTATCTAAAA                             | ATTOTATAGE  | AAGTCTACAC        |
|                          |                          | TCCCTCD DCT                  | GAATGCTTTA                             | GTATAGCGAT  | 12.0101.0.10      |
| CTCCCTCAAA               | GTATACTGAT<br>CTCATTCGAT | AAAACAAAGI                   | TTCDCDDDCD                             | CTTATAAGTT  | ATTTCTAAAG        |
| TTATCGTATT               | TAAACTCCCA               | AAAACAAAIG                   | TTCAGAAACA                             | CCAAAGCCTT  |                   |
| GGCACCTCTA               | TAAACTCCCA               | MARIIGCGAA<br>mmmacaaaaa     | TACTTTTTAC                             | AGCTCCCCTA  | AAATAGAAGA        |
| GTTAAATCAA               |                          |                              |  | TTGTAGAAAA  |                   |
| TAACAGAAGG               | GAGCCTTCAA<br>CAATATTTAG | AMACIICAII                   | CTTCACTTOGA                            | TEGEGGGGAGA | CCTTAATAAT        |
| ACTGTTAAAT               | TATTTCGAAA               | AIIIIIAGGA                   | CTADAATCAG                             | AACTGATGGT  | 0011111111        |
| CTATGCACTA               | AAGAGAATGA               | CCARTTATC                    | ידידים מידים ממממ<br>מידים מידים מממממ | CTTTTACAAT  | GGTTAATAAC        |
|                          |                          | ATTTATACGG                   | ፈተፈተፈተ ተ                               | ACTTTATTGA  | 001111111         |
| GAAAGTGAGA               | ATTATTGATA               | ATTIMIACOG                   | ACATAACACG                             | ATGCAAATTA  | TTTAATTT          |
| CGAGATGGTC               | GGATATAAAA               | MIGGIIGIAC                   | TCATCACTCT                             | TTAGAGGCAT  |                   |
|                          |                          | TAICCGIAIA                   | TANCCAAAAT                             | AATTGCTGAA  | AAAAATCCAG        |
| ATAATCAGTA               | ACCTTTGAT                | CCCCATCAAT                   | TARCOAMA                               | CGATTCAAAT  |                   |
|                          | ACCITIGGAT               | A CTCC A CTTA                | CANAGATAC                              | ATTATGTGAA  | TTGGCAATGG        |
| CCACGGAAAC               | CTAAAAAAAA               | MCIGGACIIA<br>MCNUNUUNNUUNNU | CATTCCTTTA                             | TACCACGTAG  | 11000             |
|                          | CTAAAAAAAAA              | AACCTCTTTC                   | CCATCATTCT                             | GATGGTAAAC  | CAGTTACTAA        |
| AATGCAATAT               | TGTTTTGAAA               | AMCCIGITIO                   | ATCALTCL                               | AAGCTATCGA  | 0.101111011-      |
| ATGTATAATT               | CACTGTTTTT               | CCTAACCCAA                   | ATCTAACCAT                             | AGAACATCAT  | AATGATTTGA        |
| TGGGACATCA               | TTATCGAGCT               | NTTNCCCAA                    | ACCDATTAAT                             | TTATAAAACA  |                   |
| AATTTGCACA               | CTATTCGCGA               | TATTCCTACT                   | ATCCACAACA                             |             | AGCTCAAAGA        |
| ATTTGTTACA               | TGGCGCTCAT               | TCAATCTGGC                   | GTGGATATGT                             | GGGAAACGGC  |                   |
| -                        | TGGCGCTCAL               | CTTATCATTC                   | ТААТСТТАТА                             | CATGCACCAA  | TTGATTTAAG        |
| GAGAGAAGCC               | GAAAATATTG               | TADADATA                     | TAACGAACTA                             | TCCAGAGAAA  |                   |
| TTTTTGTAAA               | ACGCGTGATG               | ADADCGGGAA                   | GAGAAATGGC                             | TGTTCGTGCA  | TATAATGTGG        |
| CAGTAGCAGA               | AAAAGAAAAG               | ΔΔΔΥΥΥΤΟΥΑΑ                  | AACCTATTAT                             | ATTTGTATTA  |                   |
| AGCGAAAACA               | AAAGGAGATGA              | GTATATTCAT                   | CCCAATCCAT                             | CAAATCATTT  | GACGATCTTA        |
| GATGGGIIAA               | ATAACGTCAG               | AGGCTTACTT                   | ACCGATAATC                             |             |                   |
| ACTGAAAIGI               | GTTAATTATA               | CATTAATTAT                   | AACTCCAGAT                             | TTTGCTAAGT  | TTTTACCGCA        |
| MCD AUTOTOMA             | GTTGTACCAG               | ATACCTTGGA                   | TATAGAGCAA                             |             |                   |
| AGTATGTTGG               | TACAGGTGTA               | GACTTGTCAA                   | AGATTATTTC                             | TTTAAAAGAG  | TATCGAAAAG        |
| AGATAGGCTT               | TATTGGTAAT               | TTGTATGCGC                   | TTTTAGGATT                             | TGTTCCGAAT  |                   |
| AGAIAGGCII               | GAATTTATCT               | ATATATTCAG                   | AGAAACGGTA                             |             | TATTATAAAA        |
| ATGCTCAATA               | GATTGTGAGA               | GTTGTTTACT                   | TTTATTTGTA                             | ATTTTAAAAG  |                   |
| TANTEC AGEC              | AGATAGGAGA               | AAAACGTTTG                   | GAAAAATGAG                             | AATAAGAATT  | AATAATTTGT        |
| TAATGCAGGC               | CATAGCGTTT               | ATGGGCATAA                   | TTATTAGTAA                             | TTCGCAAGTT  |                   |
| CTTCTACCCA               | TAGGCAAAGC               | TTCTGTGATT                   | CAGTATCTAT                             | CTTATTTAGT  | TTTGATTTTA        |
| TCTATACTTA               | ATGATTTATT               | AAAAAATAAC                   | AAACATATTG                             | TAGTTTATAA  |                   |
| TOTALAGILA               | TTGTTTCTTA               | TTATATTTT                    | ATTTACTATC                             | GGAATATGTC  | <b>AGCAAATTCT</b> |
| WCCADADDCD<br>WCCADADDCD | ACTAAAATAT               | ATTTATCAAT                   | TTCAATGATG                             | ATTATTTCAG  |                   |
| TCCIVITYCV               | GTTGCCAATA               | AGTTTGATAA                   | AAGATATTGA                             | TGATTTTAGA  | CGGATTTCAA        |
| TITINGCARC               | ATTCGCTCTT               | TTTATAACTT                   | CGATATTAGG                             | AATAAAGATG  |                   |
| AICAILIGIL               | TGTTCACGGG               | GCAGTAGAA                    | GGTATCGGTT                             | TTAGTCAGGG  | TTTTAATGGA        |
| CCATTCACCC               | ATAAGAACTT               | TTTTGGAATA                   | ACTATTTTAA                             | TGGGGTTCGT  |                   |
| COATIGACGC               | TTGGCGTATA               | AGTATGGTTC                   | CTATAAAAGA                             | ACGGATCGTT  | TTATTTTAGG        |
| ATTAACTTAC               | TTTTTGATTC               | ΤΤΑΤΤΤΟΟΙΙΟ                  | CACACGCTCA                             | GTTTATTTAA  |                   |
| ATTAGAATTG               | TTTTCTATTT               | СТТСТТОТТ                    | TTGACAAAAT                             | CAAAATAGAA  | CAAAGACAAT        |
| TACTATTGCT               | TAAATATATT               | ТССАТССТАТ                   | TTTGTGCTAT                             | TTTTTTATAC  |                   |
| GGAGTAUGUT               | GTTTTTTAAT               | AACACATACT                   | GATTCTTACG                             | CTCATCGCGT  | TAATGGTCTT        |
| TATTTCTTTG               | TTGAGTATTA               | ТАСАДАТСАТ                   | TGGTTCCATC                             | TAATGTTTGG  |                   |
| ATTAATTTTT               | TTGGCATATG               | CCCATTTAAC                   | TTTAGACTAT                             | GCTATAAGGG  | TTAGACGCGT        |
| TGCAGCGGAT               | AATGGAACGC               | TTGAAATGCC                   | CTTACTGAGT                             | ATTATGTTAA  |                   |
| TTTAGGTTGG               | AN I GOWACOC             | TTOPHATOCC                   | -11.1010101                            |             |                   |
|                          |                          |                              |  | _           |                   |

# WO-00/05378 TEVISE MARGUNS

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|             |              | //:        | 9           |            | •   |
|-------------|--------------|------------|-------------|------------|---|
| AAAATGGTTT  | TATCGGTCTG   | GTAGGGTATG | GGATTGTTTT  | ATATAAACTT | TATCGTAATG  |
| TAAGAATATT  | AAAAACAGAT   | AAAAATATAA | CAATAGGAAA  | GTCTGTATTT |   |
| ATCATTGTAG  | TCCTATCTGC   | AACAGTAGAA | AATTATATTG  | TAAATTTAAG | TTTTGTATTT  |
| ATGCCAATAT  | GTTTTTGTTT   | ATTAAATTCT | ATATCTACTA  | TGGAATCAAC |   |
| TATTAACAAA  | CAACTGCAAA   | CATAAATTGG | CAGGAATAGA  | GTTTTGAGTT | GCTATTAATT  |
| TGGTAGAGCA  | TATGTTCTAT   | AGGTGGCAAG | ATAAAGATAG  | TATTTTTAC  |   |
| ATGATGATTT  | TTATGATAGC   | AAAGCAAGTT | ACGGCATAAA  | AGGAATTAGA | GGATGGAAAA  |
| AGTCAGCATT  | ATTGTACCTA   | TTTTTAATAC | GGAAAAGTAC  | TTAAGAGAGT |   |
| GTTTAGATAG  | CATTATTTCC   | CAATCGTATA | CTAATCTAGA  | GATTCTTTTG | ATAGATGACG  |
| GTTCTTCAGA  | TTCATCAACG   | GATATATGTT | TGGAATACGC  | AGAGCAAGAT | CAAMMACCCM  |
| GGTAGAATAA  | AACTTTTCCG   | GTTACCAAAT | GGTGGTGTTT  | CAAACGCAAG | GAATTACGGI  |
| ATCAAAAATA  | GCACAGCAAA   | TTATATTATG | TTTGTAGATT  | AAAGAGAATG | አጥአ <i>ር</i> ጥ <b>ር አ</b> ጥጥጥ                     |
| TGTTGACGGC  | AACATTGTTG   | AGICCITATA | AAATTATCAA  | CARCTCACC  | HINGIGHIII  |
| GTCGGGAGGG  | TTACTTGCIA   | CITITUALGG | TARALIAICAA | GCGAGACTTA | CCAAATCAAA  |
| TGCAAAAGTG  | TCAAATTGAT   | ACCCCTATCT | TTAATAGCCC  | TTCTTCCAAA | GONNA I GANA                                      |
| ATTTTCCCAA  | TCATIATATO   | AGCGGIAICI | TTTCACACTC  | AACAGTGGTT | ACCACACCAC  |
| CTTTATAAGA  | ATAIAIAIAIA  | TTTDDDCDAT | ATAAAAAAAG  | TCCGCTATGT | noononoone  |
| TTATTATTA   | VICIWWITT TO | CCACAACAAC | TTTDCAAAGT  | ACTACAAATA | ССТТТАВАТА  |
| TAACAGAAA1  | ATTCAATTAG   | AAAATTTAGA | AGAAAAAACT  | TTTGATTTGT | 00111111111                                       |
| TURIUITIT   | ATTTGGTGGA   | CAATATGAAT | TTTCTGTTTT  | TAAAGAGACG | CTACAGTGGC  |
| 11011አለሌና፣  | TTATAGCTTA   | TTAATGTTCA | AAAATGGAGA  | TGAATCGCTT | •   |
| CCAAAGAAAT  | TGCATATATT   | TAAGTATTTA | TACAATAGGC  | ATTCTTTAGA | TACTCTAAGT  |
| ATTAAACGAA  | CGTCCTCTGT   | TTTTAAAAGA | ATATGTAAAT  | TAATTGTTGC |   |
| TAATAATTTG  | TTTAAAATTT   | TTTTAAATAC | TTTAATTAGG  | GAAGAAAAA  | ATAATGATTA  |
| ACATTTCTAT  | CATCGTCCCA   | ATTTACAATG | TTGAACAATA  | TCTATCCAAG |   |
| TGTATAAATA  | GCATTGTAAA   | TCAGACCTAC | AAACATATAG  | AGATTCTTCT | GGTGAATGAC  |
| GGTAGTACGG  | ATAATTCGGA   | AGAAATTTGT | TTAGCATATG  | CGAAGAAAGA |   |
| TAGTCGCATT  | CGTTATTTTA   | AAAAAGAGAA | CGGCGGCTA   | TCAGATGCCC | GTAATTATGG  |
| CATAAGTCGC  | GCCAAGGGTG   | ACTACTTAGC | TTTTATAGAC  | TCAGATGATT |   |
| TTATTCATTC  | GGAGTTCATC   | CAACGTTTAC | ACGAAGCAAT  | TGAGAGAGAG | AATGCCCTTG  |
| TGGCAGTTGC  | TGGTTATGAT   | AGGGTAGATG | CTTCGGGGCA  | TTTCTTAACA |   |
| GCAGAGCCGC  | TTCCTACAAA   | TCAGGCTGTT | CTGAGCGGCA  | GGAATGTTTG | TAAAAAGCTG  |
| CTAGAGGCGG  | ATGGTCATCG   | CTTTGTGGTG | GCCTGGAATA  | AACTCTATAA | > m < > > m = < m m                               |
| AAAAGAACTA  | TTTGAAGATT   | TTCGATTTGA | AAAGGGTAAG  | ATTCATGAAG | ATGAATACTT  |
| CACTTATCGC  | TTGCTCTATG   | AGTTAGAAAA | AGTTGCAATA  | GTTAAGGAGT | <b>ゕ</b> ぐゕぐヽ゚゚゚゙゚゚゙゚゚゙゚゚゙゚゚゚゙゚゚゚゚゙゚゚゚゚゚゚゚゚゚゚゚゚゚゚ |
| GCTTGTACTA  | TTATGTTGAC   | CGAGAAAATA | DACCARRCO   | TTCTAGTATG | ACTGACCATC  |
| GCTTCCATTG  | CCTACTGGAA   | TTTCAAAATG | AACGAATGGA  | CATTTTTAGC | COUNTCOUR   |
| AGTAGAGGAG  | ATAAAGAGCT   | TCATTCCTTC | AGCAAACAGC  | DADAGAGCT  | CITIGCIGIT  |
| TTGTTTTTAG  | GCAAATATAA   | TCATIGGIIG | ACAAMCAGC   | CAAAATAAGC | CACTTCCTTT  |
| TCTCCAAACG  | COMMANDAM    | TECTAGECTE | TCTTCATCTT  | AATTTTAGTG | ductionii   |
| ACTAATGAAT  | ANCECCEDAN   | CATABAATTC | AAGAAAGATT  | GAGAAGAAGT | GAAAGTAGTA  |
| COCCCTANCA  | ATCTTCTAAT   | ABATGGTTGA | AAGAAAAGGG  | GATTAAAATG |   |
| CICGGIAAGA  | ATACTACAAT   | AGCACTCTTT | GATACGATTA  | AATGTATCAT | GGTACTTTGT  |
| CULT TUTTED | CACATCTGGA   | TTGGTCTGTT | GAGCAGCGTC  | AATGGTTTAT |   |
| CTTTTTTT    | TTCGTTGACA   | TGGCTGTTCC | AATTTTTCTG  | TTGCTTTCTG | CCTATTTTCG  |
| AACGAATAAG  | TGGAATACAA   | AACAAGAGAC | GCTAAAGCTC  | AAGTTCAGCA |   |
| CTCCTATAAA  | AGAAAGTATA   | AACATGCTTT | GTCTCTATGC  | TATCGTGATG | GCTGTTAATG  |
| TTTTATTGAG  | CTATTCGAGA   | ACCATCTGAT | AGGAGTAAAG  | CCTTTTTCAG |   |
| GTTCTTCATC  | GCTCCGTTCA   | TTTGTCCTGT | GGCTACTTTC  | TGGAGAATCG | GGTCCAGGGA  |
| GTTGGGAGTT  | ACTATGTTCC   | GTTGTTGATT | CAGGTAGTTT  | TTTTATTACC |   |
| AATTTTGTAT  | GTTCTTTTCG   | AGAAAAATAA | ATGGTTGGGC  | TTGCTTACTT | GTTTTTTAGT  |
| ΔΔΔСΤΤΤΤΤΟΑ | GTGGATGCCA   | TATTTGCTAA | CATGGCTGAA  | CACGGCATAT |   |
| ATATATAGAC  | TAATATCACT   | TCGTTATCTT | TTTGTTCTAG  | GGCTTGGTTT | TTTCTTTCAA  |
| ACCAGGATGT  | GCGTTCCAAG   | GTAGATACTT | TCATTGCGAC  | CCTATTTGGG |   |
| ATTATTGGAG  | CAATTCTGAT   | TTTTGTGAAT | CATTCTATAG  | AGCCCTTCTC | CTGGTTTTAT  |
| GGTTGGAAGT  | CTACTTCCTT   | TCTATGCGTC | CCATTTGCGT  | ATGCTATGCT |   |
| ATTTTTTATG  | ATAAAGTATG   | GACAGAAGAT | TCCAGCAATA  | CTGTTGTCAA | AATTGGGAGT  |
| TGCTTCTTAT  | CATATCTACT   | TGACCCAGAT | GCTGTATTTT  | TCAGTAGTCG |   |

Change Marguns

|            |            | 8/!          | 50               |   | •                                      |
|------------|------------|--------------|------------------|---|--|
| CACCATTTTT | AGCAGTGCAA | TTTAAGGTAT   | CTTCGTTGAA       | TTTGTĞGAAC  | GGCTTGTTTA                             |
| CCTTTCTAAT |            | CCTCCCTATA   | TTTTCTACAA       | AGTGGATCTG  |  |
| TTTATGAGAG |            | ACCATAATCA   | CTCATTTCAG       | ATTAGCAGAT  | GCCATTTCGT                             |
|            | ATTCGCATGT |              |                  |   | <b>~~~</b>                             |
| TTATTAGCAG | GGAGTGGTAT | CTCTTTCCG    | ACCCCACTAT       | СССССТТТАТ  | тттсарасст                             |
| TGACGAGAGA | ACTGAACAAG | CIGITICIAN   | ACCCCAGIAI       | TTTGTTTAAT  | 1110122.001                            |
| ATATTTATTA | TCTGGTTCGA | DACCOTTACC   | CANTANANC        | ስጥር/ርጥስጥ <b>ጥጥ</b>  | TECACEGGAA                             |
|            |            | MAGGCTTACC   | AAMMCAAMCA       | CCTTCTTTTC  | TOURCOOURT                             |
| ACCCATGATT | TTTCACACGA | TIGATGIGGC   | CACAAACCEA       | TA A C C C C C C C  | አሮሮምሮሞልሞልል                             |
| AGAAAGAAGA | CATCTATGTC | AGTACGGATT   | CAGAAAIGIA       | A A TOTAL A CAT   | ACCICIAIM                              |
|            | TTGCGAATTT | GGAGTTACGA   | MAGCCIIGII       | GATTTGCGAG  | n CANCAGECA                            |
| CTTAAATTTT |            |              | TCCCCAAGGG       |   | ACAAGAGGCA                             |
| TCAATGTATT |            | AAAGAACTAT   | CTACTTATCA       | TACICCAICG  | አመአመመም <u>ሮ</u> ሞምሮ                    |
|            | GTACGCACTT |              |                  | TGAAGATTGT  | AIAIIIGIIC                             |
| TTCTGCAAGT | CACCTCACCG | TTACGGACTG   | GCGAACAGAT       | AAAAGAAGCC  | mc>>cccc                               |
|            | ACTTACAGGG | GGACTCAGAA   | AATGTTTTGC       |   | TGAAGGGCAA                             |
| GAAAGAGTGA | ATCAGTACAT |              | GTACAGGGGT       | TATAAAAAGG  |  |
| GGTTACTTAT |            |              | AGGAGAAAA        |   | TTTATATTTG                             |
| CCATACGATG |            |              | GTTAAAGATG       | GACGTTGAGA  | ~~ ~~~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ |
| GAGATAGTTT | GATGTCCGTT | GATATCATCG   |                  | AGATGTCAGG  | GAGCAACTGC                             |
| AGCAGCATGT |            |              | GAGCGTTCAT       | TTGATCTATA  |  |
| TTCTTTGATA | GCTAGATCAA |              |                  |   | GCTATGACGA                             |
| GGTGATCATT |            |              | CGGTCATTTT       | TTAAATAAAC  |  |
|            | CTATTCTCTT |              | GTTATAATTT       | TTTCAAGGAT  | AAAAGAGTGT                             |
|            | GTCAATTCAA |              | GGAAAAGACT       | CTTTTATCAA  |  |
| TGGTATTTTA | AACCAACATA |              |                  | ATTGTCAATC  | CATTGAGGTC                             |
| AATGATCTGT | CGCTCGTACA | ATTTGACTAG   | GCTTATAAAC       |   |  |
| AGTTCCGAGA | AAGCAATTAT | TTGATCAAGC   | ATCGCCAGAG       | AAGGTGCAAG  | CGCTGCTGCA                             |
|            | GCAAGGGCGA |              | TGAAGAGTCT       | TCTCAAAAAC  |  |
| GATTGCTATT | ATTGACCCAG | CCCTTGTCTT   | GGGATTATCA       |   | GAGAGTTGTT                             |
| GGAGATTTAT | GTAGCAGGTC |              | TCGGGAAGAC       | TATACAATCT  | •                                      |
|            |            |              |                  | TCTGGGTAAG  | GCTGTGGTGC                             |
|            | AGGTATTCCG |              |                  | AGGTAATATC  |  |
|            | TCGGTATGAC |              |                  | ATTTTTTAAA  | TTGTTTTGAA                             |
| GAGAAAGTGT | ATTTAAAGGA | CACTTTTCCT   |                  | AAAATGATAT  |  |
| TTTGCGTGAG | GGGATAGAAT | AGGAGGATTC   | ATGTCTAAAA       | AATCAATAGT  | TGTCTCAGGT                             |
| CTCGTCTATA | CGATTGGAAC | CATCCTCGTT   | CAGGGATTAG       | CCTTCATTAC  |  |
| CCTCCCCATC | TATACTCGTG | TCATTTCTCA   | GGAAGTATAT       | GGGCAGTTTA  | GCTTGTATAA                             |
| TTCGTGGGTG | GGGCTAGTTG |              | CGGTCTACAG       | TTAGGTGGGG  |  |
| CTTTTGGCCC | GGGATGGGTA | CACTTCCGCG   | AGAAATTTGA       | TGATTTCGTA  | TCCACCTTGA                             |
| TGGTCTCTTC | TATCGCTTTC | TTTTTACCAA   | TTTTTGGGCT       | ATCTTTTCTC  |  |
| CTCAGTCAGC | CCCTATCGCT | CCTATTTGGT   | TTGCCTGATT       | GGGTCGTTCC  | GCTTTACTTT                             |
| TTGCAAAGTT | TTATGAGTGT | TGTGCAAGGA   | TTTTTTACGA       | CCTATTTAGT  |  |
| GCAGCGGCAG | CAGTCCATGT | GGACTTTACT   | CCTATCGGTA       | CTGAGCGCTG  | TTATCAACAC                             |
| TCCTTTATCT | TTATTTCTCA | TCTTTTCGAT   | GGAGAATGAT       | TTCATCGCTC  |  |
| GTGTAATGGC | AAACTCGGCA | ACGACTGGTG   | TTTTTGCTTG       | TGTGTCCTTG  | TTGTTTTTCT                             |
| ATAAGAAGAT | TGGGCTTCAT | TTTCGAAAGG   | ACTATCTTCG       | GTATGGTTTA  |  |
| AGTATATCGA | TTCCTCTTAT | TTTTCATGGA   | TTAGGTCATA       | ATGTACTCAA  | TCAATTTGAC                             |
| ACAATCATCC | TCGGCAAGAT | GCTAACACTG   | TCAGATGTAG       | CCCTATACAG  |  |
| TTTCCCCTAC | ACACTTGCGT | CTATCTTACA   | AATTGTGTTT       | TCGAGCTTGA  | ATACGGTATG                             |
| CTCTCCCTCC | TATTTTGAGA | AAAAGAGAGG   | TGCAGATAAA       | GATTTGCTCA  |  |
| GIGICCGIGG | TTACTATCTG | GCGATTGGCC   | TGTTTGTGAC       | TTTTGGATTT  | CTAACAATTT                             |
| GITATGICCG | AGCGATGTTG | ттасстссат   | CTGAGTATCG       | TTTCAGTATG  |  |
| ACCCTGAATT | CCATGATTAT | TCTCCCCCTC   | TTCTTTGTAT       | TTCTTTATAG  | TTTTCCAGCC                             |
| GGATTTATTC | TTTATAGTGG | TOTOGGGTG    | ጥጥጥጥርርርልል        | TTGGTACTTT  |  |
| AATATCCAGT | GTACTAAATA | TATA TACAMAG |                  | ATACCGACAA  | AGAATTTATG                             |
| TATAGCAGGT | GCAACGACTG | TITCCGTCCA   | CTITOIIII        | CTCTTCCATT  |  |
| GTGCTGCTTT | TAAGAAAAAG | CITCCIAICI   | OTIGITOCIA       | CATTTCALL   | <b>ΤΤΤΩΤΤΆΔΩ</b> Ω                     |
| ATTTTGTTGC | TAAGAAAAAG | TAIGUTTAUG   | TIGHAGIIGC       | <b>Δ</b> ጥጥተርጥርርርጥ  | 111011111100                           |
| TAATTGCTCT | TGTTGTCGTC | TATACAGGCT   | 1 GW 1 GW CW G 1 | TITIGIOGI   | СФДСДФФФФФФ                            |
| TCAATCTGGA | TTCGTTGGTC | ACTAGGAATA   | TTC ACCCA A A    | A D C C C T C T A A C C C T C T A A C C C T A A C C C T A A C C T A A C C T A A C C T A A C C T A A C C T A C | CINCALLLE                              |
| AGAAAGGAAT | TAACAGTTGC | CCTCAATACA   | 1 I CAGGGAAA     | MUCGGICIAM  |  |
|            |            |              |                  |   |  |

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|            |            | 9/:        | 50          |              |                         |
|------------|------------|------------|-------------|--------------|-------------------------|
| ATAAGGGCAC | CTCTATAAAC |            | GCGAATTTGG  | AGTTACGAAA   | GCCTTGTTAA              |
| ATCAAACATT |            | GAAAATTAGT |             | CCCCATATAA   |                         |
|            | AATGAGAGGT |            | ATTGACCATC  | ACTGCCATCT   | ACCCAAAGTT              |
|            | CTACCATGAA |            |             |              |                         |
|            |            |            |             | AAGTCATGAG   | GTAAATCTCC              |
|            | GGTAGAAAAG |            |             | CTATTGAAGA   |                         |
|            | GAGCAGGCTT |            |             | ACTATTCGAA   | AGAAATCTAG              |
| GGCTATTTTT |            |            |             | TCTTTATTAG   |                         |
|            | CTACTTAAGG |            |             | AGACTTTATG   | GGAGCGATTT              |
|            | TTAGAAAGGA |            | TTTATATTAT  | TGCAGAAATT   |                         |
|            | ACAACGGTGA |            | GCACGGAAAA  | TGGTAGAAGT   | TGCCGTTGAT              |
|            | ATGCCGTTAA |            |             | ATTTGTTGAT   |                         |
| TTCAAAATAC | GCACCAAAGG | CCGAATACCA | AAAAATTACA  | ACAGGAGAGT   | CAGATTCTCA              |
|            | ACTCGTCGTT |            |             |              |                         |
| TGCGTGATTA | CTGTCTTGAA | AAGGGAGTTG | ATGTGTTTTC  | GACACCTTTT   | GATGAGGAAT              |
|            | CTTGATTAGC |            |             |              |                         |
| GGTGAGATTA | CCAATCTTCC | CTATTTGGAA | AAAATTGGTC  | GTCAAGCTAA   | GAAAGTTATT              |
|            | GTATGGCTGT |            | ATTCATCAAG  |              |                         |
|            | AATGGAACGA |            | GATTTTGCAT  | TGTACAACCG   | AGTATCCAAC              |
|            | GCTTTGAATT |            | GCATACCTTG  |              |                         |
|            | AACAATTGGC |            | ATAGTGTTGG  | TTCAGAAGTA   | CCCATCGCTG              |
|            | GGGAGCTGAA |            | AGCACTTTAC  | TCTGGACAAT   |                         |
|            | GACCAGATCA |            |             | ATATCTTAGC   | AGCCTTGGTA              |
|            | GGATAGTGGA |            | GGTAAATTTG  | AAAAAGAGCC   |                         |
|            | GAAGTACGAA |            | AGCTAGAAAA  | TCTATTGTTG   | CCAAAAAAGC              |
|            | GGCGAAGTCT |            | AAACATCACT  | GTCAAAAGAC   |                         |
| CAGGAAATGG | AATTTCGCCA | ATGGAATGGT | ACAAAGTCTT  | GGGGCAGGTG   | AGTGAGCAGG              |
|            | AGACCAAAAT |            | GTGCTTTTGA  | AAATCAAATG   |                         |
| TAAGCGGAGT | AAGGATGAAA | AAAATTTGTT |             | CTCTCGTGCC   | GAATATGGGA              |
| TTATGCGTCG | CTTATTGAGC | TATCTACAGG | ATGATCCAGA  | AATGGAGCTG   |                         |
| GATCTTGTAG | TGACAGCCAT | GCATCTAGAA | GAAAAATATG  | GGATGACGGT   | CAAAGACATC              |
| GAAGCGGACA | AGCGTAGGAT |            | ATTCCATTGC  |              |                         |
|            | CAGACAATCG |            |             | ACAGAGCAAC   | TCACGGTTCT              |
|            | GTCCAGTATG |            | GATTCTGGGG  |              |                         |
| AGATGCTACC | AGTTGCCAAT | GCTGCGTTGC | TTTATAATAT  | TCCTATTTGC   | CATATTCATG              |
|            | AACCATGGGA |            |             |              |                         |
|            | GTCACCTTCA |            |             | TTAGAAATCG   | TGTCATTCAA              |
|            | ATCCAACCAT |            |             | GGGTGTTGAA   |                         |
|            | AACAAGACTT |            | GAAGAGTTGG  |              | TGGAATTGAT              |
|            |            | TGTACTCTTT |             |              | 1 macm1 ccc1            |
| TAACACAGCC | GAAGAACAAA | CGCAGGCCTT | ATTAGATGCT  | CTAAAAGAAG   | ATGGTAGCCA              |
| GTGTTTGATA | ATTGGATCCA | ATTCGGATAC | ACATGCCGAT  | AAGATAATGG   | MCCCMMCCA A             |
| AATTGATGCA | TGAATTTGTA | AAACAAGACT | CTGATTCTTA  | CATCTTTACT   | TUGUTTUCAA              |
| CTCGTTATTA | CCATTCCTTG | GTCAAGCATT | CACAAGGTTT  | AATAGGGAAT   | <b></b>                 |
| TCTTCGTCAG | GTTTGATTGA | AGTGCCCTCA | TTACAGGTTC  | CGACCTTAAA   | IAIIGGAAAI              |
| CGCCAATTTG | GACGTTTGTC | AGGACCGAGT | GTGGTACATG  | COCAMACATO   | መመስ ርር እ አጥርር           |
| TAAGGAAGCG | ATTGTTGGTG | GTTTGGGGCA | ATTACGTGAT  | GTGATAGATT   | TIACCAATCC              |
| ATTTGAACAA | CCTGATTCTG | CTTTACAAGG | TTATCGAGCI  | MICAMOGRAI   | አርአ አ አ <b>ር</b> ጥጥጥር   |
| TTTTATCTGT | ACAGGCCTCA | ACCATGAAAG | AGTITIATGA  | TAGATAGGG    | AGAAAGIIIG              |
| ATGAAAAAAG | TAGCCTTTCT | AGGAGCGGGT | ACCITITICAG | MUDGIGICCI   | <u>አአርርርአሞርአር</u>       |
| TCCTTGGTTG | GATAGAACTC | GATATGAACT | CMTTGGATAT  | TTTGAAGATA   | NACCONTCAG              |
| TGACTATCGT | GGCTATCCTG | TATTTGGTCC | CTTGCAAGAT  | DICCIAACCI   | CTCNACCCCA              |
| ATTTGGATGA | TGGAAAAGTA | GATGUTGTUT | AMMAMOAMOO  | AGGTGACAAT   | GICMMOCOCH              |
| AGGAAATCTT | TGACTTGCTT | GCCAAAGATC | ATTATGATGC  | TITGITCAAC   | NCCCCMMMMC              |
| ATCATTAGCG | AGCAAGCCAA | TATTTTTCC  | CCAGATAGTA  | TCAAGGGACG   | MOGGGIIIIC              |
| ATAGGTTTTT | CAAGTTTTGT | AGGAGCCGAT | TCCTATGTCT  | CTCCACCCCC   | አጥጥሮመን አሮአማ             |
| TATCATCAAT | ACGGGTGCCA | TTGTGGAACA | TUATACCACG  | GTGGAGGCCC   | MIIGIAACAT              |
| TACTCCAGGA | GTGACCATAA | ATGGCTTGTG | CUSTATUGGA  | TOTAL COMPAN | ארא ארא <b>יי</b> ייייר |
| ATATTGGAAG | TGGTTCAACA | GTGATTCAAT | GTATUGAGAT  | TGCACCTAT    | ACAMCA1 166             |
| GGGCAGGGAC | AGTTGTTTTG | AAATCGTTGA | CGGAGTCAGG  | GACCIAIGTT   |                         |

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| •           |               | 10.        | /59           |            | *                          |
|-------------|---------------|------------|---------------|------------|----------------------------|
| CCTCTACCTG  | CTAGAAAGAT    | TAAATAGGTG | AATTGATGGA    | ACCAATTTGT | CTGATTCCTG                 |
|             | ATCAAAAGGT    | TTACCAAATA | AAAACATGTT    | ATTTTTAGAT |                            |
|             |               |            |               | AGTCTGGATG | TTTTAAGAAA                 |
| GGTGTACCGA  | ATGTCAGTAC    |            | GTTTACAAGG    |            |                            |
| GAAAATATAT  |               |            |               | GCGACAGATT | TTACAACCTC                 |
| AACAACTGGG  | GTTCAAGTCC    |            | TTTTTCTGAT    |            | 1111011110010              |
| TTTTCAACTG  |               |            |               |            | CACCCATCC                  |
| TTGTTCTCCT  | GCAAGTTACG    |            |               | ACATGTCAAG | GAGGCGAIGG                 |
| AGTTATATGG  |               | GCTGACCACG |               | TACCAAAGTC | ~~~ ~~ ~~~                 |
| GATAAGTCTC  | CAACATTGTT    |            | GACGAAAACG    |            | GGATATTGCA                 |
| GGATTAGGTG  | GCAGTTATCG    |            | GAGAAAACAC    | TCTACTATCC |                            |
| TAATGGAGCG  | ATTTATATTT    | CTTCTAAGCA | GGCTTATTTA    | GCGGATAAAA | CTTATTTTTC                 |
| TGAAAAAAACA | GCGGCCTATG    | TGATGACGAA | GGAAGATTCG    | ATTGATGTAG |                            |
| ATGATCACTT  | TGATTTTACT    | GGTGTTATTG | GTCGAATTTA    | CTTTGATTAC | CAGCGTCGTG                 |
|             | CAAACCATTT    | TATAAAAGAG | AGTTAAAGCG    | TTTATGTGAG |                            |
|             | ATGATAGTCT    | TGTGATTGGC | GATAGTCGTC    | TGTTAGCCTT | GTTACTGGAT                 |
| CCTTTCGATA  | ATATCAGCAT    | CGGTGGGATG | ACAGCTTCGA    | CAGCACTTGA |                            |
| AAACCAAGGT  | CTCTTTTTGG    | CTACTCCGAT | AAAGAAAGTT    | TTGCTTTCTC | TTGGTGTGAA                 |
| MCAMPTCATT  | ACTGACTATC    | CCTTGCATAT | GATTGAGGAT    | ACTATTCGCC |                            |
|             | AAGTCTTGTT    |            | AGCAGGTTTT    | TGTGACGACG | ATTGCCTACA                 |
|             |               | TCCAATGAAG | AAATTGTGCA    |            |                            |
| CGCTGTTTCG  | AGTCAGCAAG    |            |               | <b>+</b> · | TGAAGTTGTT                 |
| GTTATTGTTC  | CGATGCTTGA    | CTATCACTAT | ACCAATGATG    |            |                            |
| GAAAAAGAGG  | CCALGCIIGA    | CIMICAGIAI | CCTCATUATO    | ACAAGTTTGA | САВСАТАВТТ                 |
| CAATCAGATT  | - COMPARENCE  | GIGIGAAICA | GCIGATITIG    | TGTGTTTTAG | 0.1.0                      |
| TGGTGATAGA  | AGCTATTTCA    | DIGGCIAGAC | ACCCUMCACA    | ATTCACAACC | <b>Δ</b> ጥጥ <b>Δ</b> ርጥጥርጥ |
|             |               | AGAGGATGCT | AGCCIIGAGA    | ATTGACAACC | Allinoiloi                 |
| TTTAATTATA  |               | TCTAAAAACT |               |            | <b>CNCCNTCNNN</b>          |
| AGATAATAGA  | ATAAAAAGTA    | ATGAGGAGAG | CTGTCATGCA    |            | GACGAIGAAA                 |
|             | AAAACTATCA    | GAGAAAGGCA | ATCCCTTAGA    | ACGITIGGAL | መርረሞክ አ አርአጥ               |
| GCCGTTATGG  |               |            |               | AGTTATTCAG | ICGIAAAGAI                 |
| AAAGTCATCA  | GTCGTGGCGG    |            | CTAGACTATC    | TCATGATGTT |                            |
| CAAAGCGCTC  |               |            |               | GATGCCATGG | AATATCAACT                 |
| GCTGGATCGT  | ATATCTTTTC    |            | TGGTTGTCAT    |            |                            |
| TTCCCGATGC  | GAAAACTATC    |            |               | AACCAAGTCA | GGTCGTGAAA                 |
| AGGAGTTGTT  | CGATTTGTTC    |            | TCACAGATGA    | AGGGGTGATT |                            |
| GCCCATTCAG  | GTCAGATTGT    | GGATGCTACC | TTTGTCGAAT    | GCCCTAAACA | ACGCAATTCA                 |
|             | ATCAGAAAAT    | CAAAACTTAT | CGAAAATTAT    | GAGGTCACAA |                            |
|             | ACACGACTCC    | AATGTCCTAG | CTCCTCTTTG    | TGATGCCAAT | GAAGCGGTTT                 |
| TTGATGACAG  | TGCTTATGTT    |            | TACCAGAAGG    | TTGTCGCCAC |                            |
| CACACGATTC  | GTCGTGCTTT    | TAGAAATAAA | CCGTTGACTG    | AGACTGATAA | GGTCATTAAT                 |
| CCACATATTA  | CCAAAGTCCG    | TTGTCGCGTT | GAGCATGGTT    | TTGGCTTCAT |                            |
| TCNANCTAAC  | ATGAAAGGTA    | ACATCTGTCG | AGCAATTGGG    | AAGGCACGAG | CTGAAACCAA                 |
| TGAAACIAAC  | ACCAACCTGC    | TCTACAATAT | CTGTCGTTTT    | GAGCAAATCA |                            |
| BACCACTCC   | ATTACCATCC    | GTGGGCTTAG | TGCGCCCAAA    | AAATAGGAAA | ATAAGCAAAA                 |
| AACGACIGGG  | CAAAAACTAG    | ΨΨΨΟΨΟΑΛΑΑ | TAAAAAAACG    | GCTCTTTGTC |                            |
| AGAGGC1GGG  | CAMAMCIAG     | AACCTAACAC | CTACACACGA    | CGAAATTCGT | TCTCTCATTT                 |
| AACTGTAGTG  | D D CCCUDA CC | CCCTAATAAC | A A C CTATCTA | TCCAATCACA |                            |
| TTGATGTTTA  | AAGCGTAACC    | GCCIMMIMAC | ANGUINICIA    | ATCTATGATA | ТААТСТАТТТ                 |
| CATTCCTCCA  | TTATATAGTT    | MAATGAAACA | CMMCMCCCCAM   | ACTCTCACAA |                            |
| ATGGCATATT  | CATTAGATTT    | TCGTAAAAA  | GIICICGCAI    | ACTGTGAGAA | СФВФСФВФСВ                 |
| AACCGGCAGT  | ATTACTGAAG    | CATCAGCTAT | TTTCCAAGTT    | TCACGTAACA | CIMICINION                 |
| ATGGCTAAAA  | TTAAAAGAGA    | AAACCGGCGA | GCTTCATCAC    | CAAGTTAAAG | A CTICA TICCA C            |
| GAACCAAGCC  | AAGAAAAGTG    | GATAGAGATA | AATTAAAGAA    | TTATCTTGAA | ACTUATUCAG                 |
| ATGCTTATTT  | GACTGAAATA    | GCTTCTGAAT | TTGACTGTCA    | TCCAACAGCT |                            |
| ATTCATTACC  | CCCTCAAAGC    | TATGGGATAT | ACTCGAAAAA    | AAAGAGCTGT | ACCTACTATG                 |
| AACAACACCC  | TCAAAAAGTA    | GAACTGTTCC | TTAAAGAATT    | GAATAACTTA |                            |
| ACCCACTTGA  | CTCCTGTTTA    | TATTGACGAG | ACAGGGTTTG    | AGACATATTT | TCATCGAAAA                 |
| ጥአጥርርጥርርርጥ  | CTTTGAAAGG    | TCAGTTGATA | AAAGGTAAGG    | TCTCTGGAAG |                            |
| מאכאיזארראר | CGGATATCTT    | TAGTAGCAGG | TCTCATAAAT    | GGTGCGCTTA | TAGCCCCGAT                 |
| CACAMACAAA  | GATACTATGA    | CGAGTGGCTT | TTTCGAAGCT    | T          |                            |
| GWCWIWCWUW  |               |            |               |            |                            |

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11/59

SLDIDHMMEVMEASKSAAGSACPSPQAYQAAFEGAENIIVVTITGGLSGSFNAARVARDM YIEEHPNVNIHLIDSLSASGEMDLLVHQINRLISAGLDFPQVVEAITHYREHSKLLFVLA KVDNLVKNGRLSKLVGTVVGLLNIRMVGEASAEGKLELLQKARGHKKSVTAAFEEMKKAG YDGGRIVMAHRNNAKFFQQFSELVKASFPTAVIDEVATSGLCSFYAEEGGLLMGYEVKA

Fig. 3 cont.

ORF2Z

SEQ. ID. NO. 10

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12/59
MKKYQVIIQDILTGIEEHRFKRGEKLPSIRQLREQYHCSKDTVQKAMLELKYQNKIYAVE
KSGYYILEDRDFQDHTCRAQSYRLSRITYEDFRICLKESLIGRENYLFNYYHQQEGLAEL
ISSVQSLLMDYHVYTKKDQLVITAGSQQALYILTQMETLAGKTEILIENPTYSRMIELIR
HQGIPYQTIERNLDGIDLEELESIFQTGKIKFFYTIPRLHNPLGSTYDIATKTAIVKLAK
QYDVYIIEDDYLADFDSSHSLPLHYLDTDNRVIYIKSFTPTLFPALRIGAISLPNQLRDI
FIKHKSLIDYDTNLIMQKALSLYIDNGMFARNTQHLHHIYHAQWNKIKDCLEKYALNIPY
RIPKGSVTFQLSKGILSPSIQHMFGKCYYFSGQKADFLQIFFEQDFADKLEQFVRYLNE

Fig. 3 cont.

ORF2Y

SEQ. ID. NO. 53

verise mayons



MKIIIPNAKEVNTNLENASFYLLSDRSKPVLDAISQFDVKKMAAFYKLNEAKAELEADRW YRIRTGQAKTYPAWQLYDGLMYRYMDRRGIDSKEENYLRDHVRVATALYGLIHPFEFISP HRLDFQGSLKIGNQSLKQYWRPYYDQEVGDDELILSLASSEFEQVFSPQIQKRLVKILFM EEKAGQLKVHSTISKKGRGRLLSWLAKNNIQELSDIQDFKVDGFEYCTSESTANQLTFXR SIKM

Fig. 3 cont.

ORF2X

SEQ. ID. NO. 11

change margins

1

14/59

MKKRSGRSKSSKFKLVNFALLGLYSITLCLFLVTMYRYNILDFRYLNYIVTLLLVGVAVL AGLLMWRKKARIFTALLLVFSLVITSVGIYGMQEVVKFSTRLNSNSTFSEYEMSILVPAN SDITDVRQLTSILAPAEYDQDNITALLDDISKMESTQLATSPGTSYLTAYQSMLNGESQA MVFNGVFTNILENEDPGFSSKVKKIYSFKVTQTVETATKQVSGDSFNIYISGIDAYGPIS TVSRSDVNIIMTVNRATHKILLTTTPRDSYVAFADGGQNQYDKLTHAGIYGVNASVHTLE NFYGIDISNYVRLNFISFLQLIDLVGGIDVYNDQEFTSLHGNYHFPVGQVHLNSDQALGF VRERYSLTGGDNDRGKNQEKVIAALIKKMSTPENLKNYQAILSGLEGSIQTDLSLETIMS LVNTQLESGTQFTVESQALTGTGRSDLSSYAMPGSQLYMMEINQDSLEQSKAAIQSVLVE K

Fig. 3 cont.

CPS2A

SEQ. ID. NO. 12

change mayins



15/59 MNNQEVNAIEIDVLFLLKTIWRKKFLILLTAVLTAGLAFVYSSFLVTPQYDSTTRIYVVS QNVEAGAGLTNQELQAGTYLAKDYREIILSQDVLTQVATELNLKESLKEKISVSIPVDTR IVSISVRDADPNEAARIANSLRTFAVQKVVEVTKVSDVTTLEEAVPAEEPTTPNTKRNIL LGLLAGGILATGLVLVMEVLDDRVKRPQDIEEVMGLTLLGIVPDSKKLK

Fig. 3 cont.

CPS2B

SEQ. ID. NO. 13

revised to satisfy margin requirements

MAMLEIARTKREGVNKTEEYFNAIRTNIQLSGADIKVVGITSVKSNEGKSTTAASLAIAY  ${\tt ARSGYKTVLVDADIRNSVMPGFFKPITKITGLTDYLAGTTDLSQGLCDTDIPNLTVIESG}$ KVSPNPTALLQSKNFENLLATLRRYYDYVIVDCPPLGLVIDAAIIAQKCDAMVAVVEAGN VKCSSLKKVKEQLEQTGTPFLGVILNKYDIATEKYSEYGNYGKKA

Fig. 3 cont.

CPS2C

SEQ. ID. NO. 14

revised requirements

1

17/59

MIDIHSHIIFGVDDGPKTIEESLSLISEAYRQGVRYIVATSHRRKGMFETPEKIIMINFL QLKEAVAEVYPEIRLCYGAELYYSKDILSKLEKKKVPTLNGSCYILLEFSTDTPWKEIQE AVNEMTLLGLTPVLAHIERYDALAFQSERVEKLIDKGCYTQVNSNHVLKPALIGERAKEF KKRTRYFLEQDLVHCVASDMHNLYSRPPFMREAYQLVKKEYGEDRAKALFKKNPLLILKN QVQ

Fig. 3 cont.

CPS2D

SEQ. ID. NO. 15

revised to sutisfy margin requirements 4

18/59

MNIEIGYRQTKLALFDMIAVTISAILTSHIPNADLNRSGIFIIMMVHYFAFFISRMPVEF EYRGNLIEFEKTFNYSIIFVIFLMAVSFMLENNFALSRRGAVYFTLINFVLVYLFNVIIK QFKDSFLFSTTYQKKTILITTAELWENMQVLFESDILFQKNLVALVILGTEIDKINLPLP LYYSVEEAIGFSTREVVDYVFINLPSEYFDLKQLVSDFELLGIDVGVDINSFGFTVLKNK KIQMLGDHSIVTFSTNFYKPSHIWMKRLLDILGAVVGLIISGIVSILLIPIIRRDGGPAI FAQKRVGQNGRIFTFYKFRSMFVDAEVRKKELMAQNQMQGGMFKMDNDPRITPIGHFIRK TSLDELPQFYNVLIGDMSLVGTRPPTVDEFEKYTPSQKRRLSFKPGITGLWQVSGRSDIT DFNEVVRLDLTYIDNWTIWSDIKILLKTVKVVLLREGGQ

Fig. 3 cont.

CPS2E

SEQ. ID. NO. 16

madified to comply with margin requirements

MRTVYIIGSKGIPAKYGGFETFVEKLTEYQKDKSINYFVACTRENSAKSDITGEVFEHNG ATCFNIDVPNIGSAKAILYDIMALKKSIEIAKDRNDTSPIFYILACRIGPFIYLFKKQIE SIGGQLFVNPDGHEWLREKWSYPVRQYWKFSESLMLKYADLLICDSKNIEKYIHEDYRKY APETSYIAYGTDLDKSRLSPTDSVVREWYKEKEISENDYYLVVGRFVPENNYEVMIREFM KSYSRKDFVLITNVEHNSFYEKLKKETGFDKDKRIKFVGTVYNQELLKYIRENAFAYFHG HEVGGTNPSLLEALSSTKLNLLLDVGFNREVGEEGAKYWNKDNLHRVIDSCEQLSQEQIN DMDSLSTKQVKERFSWDFIVDEYEKLFKG

Fig. 3 cont.

CPS2F

SEQ. ID. NO. 17

modified to comply with margin requirements

MKKILYLHAGAELYGADKVLLELIKGLDKNEFEAHVILPNDGVLVPALREVGAQVEVINY PILRRKYFNPKGIFDYFISYHHYSKQIAQYAIENKVDIIHNNTTAVLEGIYLKRKLKLPL LWHVHEIIVKPKFISDSINFLMGRFADKIVTVSQAVANHIKQSPHIKDDQISVIYNGVDN KVFYQSDARSVRERFDIDEEALVIGMVGRVNAWKGQGDFLEAVAPILEQNPKAIAFIAGS AFEGEEWRVVELEKKISQLKVSSQVXRMDYYANTTELYNMFDIFVLPSTNPDPLPTVVLK AMACGKPVVGYRHGGVCEMVKEGVNGFLVTPNSPLNLSKVILQLSENINLRKKIGNNSIE ROKEHFSLKSYVKNFSKVYTSLKVY

Fig. 3 cont.

CPS2G

SEQ. ID. NO. 18

modified to comply uith margin requirement

MKIISFTMVNNESEIIESFIRYNYNFIDEMVIIDNGCTDNTMQIIFNLIKEGYKISVYDE SLEAYNQYRLDNKYLTKIIAEKNPDLIIPLDADEFLTADSNPRKLLEQLDLEKIHYVNWQ WFVMTKKDDINDSFIPRRMQYCFEKPVWHHSDGKPVTKCIISAKYYKKMNLKLSMGHHTV FGNPNVRIEHHNDLKFAHYRAISQEQLIYKTICYTIRDIATMENNIETAQRTNQMALIES GVDMWETAREASYSGYDCNVIHAPIDLSFCKENIVIKYNELSRETVAERVMKTGREMAVR AYNVERKQKEKKFLKPIIFVLDGLKGDEYIHPNPSNHLTILTEMYNVRGLLTDNHQIKFL KVNYRLIITPDFAKFLPHEFIVVPDTXDIEQVKSQYVGTGVDLSKIISLKEYRKEIGFIG NLYALLGFVPNMLNRIYLYIQRNGIANTIIKIKSRL.

Fig. 3 cont.

CPS2H

SEQ. ID. NO. 19

modified to comply with margin requirement



MQADRRKTFGKMRIRINNLFFVAIAFMGIIISNSQVVLAIGKASVIQYLSYLVLILCIVN DLLKNNKHIVVYKLGYLFLIIFLFTIGICQQILPITTKIYLSISMMIISVLATLPISLIK DIDDFRRISNHLLFALFITSILGIKMGATMFTGAVEGIGFSQGFNGGLTHKNFFGITILM GFVLTYLAYKYGSYKRTDRFILGLELFLILISNTRSVYLILLLFLFLVNLDKIKIEQRQW STLKYISMLFCAIFLYYFFGFLITHSDSYAHRVNGLINFFEYYRNDWFHLMFGAADLAYG DLTLDYAIRVRRVLGWNGTLEMPLLSIMLKNGFIGLVGYGIVLYKLYRNVRILKTDNIKT IGKSVFIIVVLSATVENYIVNLSFVFMPICFCLLNSISTMESTINKQLQT

Fig. 3 cont.

CPS2I

SEQ. ID. NO. 20

modified to comply with margin requirement

MEKVSIIVPIFNTEKYLRECLDSIISQSYTNLEILLIDDGSSDSSTDICLEYAEQDGRIK LFRLPNGGVSNARNYGIKNSTANYIMFVDSDDIVDGNIVESLYTCLKENDSDLSGGLLAT FDGNYQESELQKCQIDLEEIKEVRDLGNENFPNHYMSGIFNSPCCKLYKNIYINQGFDTE QWLGEDLLFNLNYLKNIKKVRYVNRNLYFARRSLQSTTNTFKYDVFIQLENLEEKTFDLF VKIFGGQYEFSVFKETLQWHIIYYSLLMFKNGDESLPKKLHIFKYLYNRHSLDTLSIKRT SSVFKRICKLIVANNLFKIFLNTLIREEKNND

Fig. 3 cont.

CPS2J

SEQ. ID. NO. 21

modified comply with margin requirement

WO-00/05378

PCTANL99/00460

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24/59

MINISIVPI YNVEQYLSKC INSIVNQTYK HIEILLVNDG STDNSEEICL AYAKKDSRIR YFKKENGGLS DARNYGISRA KGDYLAFIDS DDFIHSEFIQ RLHEAIEREN ALVAVAGYDR VDASGHFLTA EPLPTNQAVL SGRNVCKKLL EADGHRFVVA WNKLYKKELF EDFRFEKGKI HEDEYFTYRL LYELEKVAIV KECLYYYVDR ENSIITSSMT DHRFHCLLEF QNERMDFYES RGDKELLLEC YRSFLAFAVL FLGKYNHWLS KQQKKLLQTL FRIVYKQLKQ NKRLALLMNA YYLVGCLHLN FSVFLKTGKD KIQERLRRSE SSTR

Fig. 3 cont.

CPS2K

SEQ. ID. NO. 22

modification to comply with margin requirement WO-00/05378

PCT/NL99/00460

25/59

| MSKKSIVVSG   | LVYTIGTILV | QGLAFITLPI | YTRVISQEVY | GQFSLYNSWV | GLVGLFIGLQ |
|--------------|------------|------------|------------|------------|------------|
| LCCA FCPGWV  | HFREKFDDFV | STLMVSSIAF | FLPIFGLSFL | LSQPLSLLFG |            |
| I.PDWVVPLIF  | LQSLMIVVQG | FFTTYLVQRQ | QSMWTLPLSV | LSAVINTALS | LFLTFPMEND |
| FTARVMANPA   | TTGVLACVSX | WFSQKKNGLH | FRKDYLRYGL | SISIPLIFHG |            |
| LGHNVLNOFD   | RIMLGKMLTL | SDVALYSFGY | TLASILQIVF | SSLNTVWCPW | YFEKKRGADK |
| DI.I.SYVRYYI | AIGLFVTFGF | LTIYPELAML | LGGSEYRFSM | GFIPMIIVGV |            |
| FFVFLYSFPA   | NIQFYSGNTK | FLPIGTFIAG | VLNISVHFVL | IPTKNLWCCF | ATTASYLLLL |
| VLHYFVAKKK   | YAYDEVAIST | FVKVIALVVV | YTGLMTVFVG | SIWIRWSLGI |            |
|              | RKELTVALNT |            |            |            |            |
|              |            |            |            |            |            |

Fig. 3 cont.

CPS20

SEQ. ID. NO. 23

modified to comply with margin requirement

-WO-00/05378

PCT/NL99/00460

26/59

MVYIIAEIGC NHNGDVHLAR KMVEVAVDCG VDAVKFQTFK ADLLISKYAP KAEYQKITTG ESDSQLEMTR RLELSFEEYL DLRDYCLEKG VDVFSTPFDE ESLDFLISTD MPVYKIPSGE ITNLPYLEKI GRQAKKVILS TGMAVMDEIH QAVKILQENG TTDISILHCT TEYPTPYPAL NLNVLHTLKK EFFNLTIGYS DHSVGSEVPI AAAAMGAELI EKHFTLDNEM EGPDHKASAT PDILAALVKG VRIVEQSLGK FEKEPEEVEV RNKIVARKSI VAKKAIAKGE VFTEENITVK RPGNGISPME WYKVLGQVSE QDFEEDQNIC HSAFENQM

Fig. 3 cont.

CPS2P

SEQ. ID. NO. 24

Change margins

MKKICFVTGS RAEYGIMRRL LSYLQDDPEM ELDLVVTAMH LEEKYGMTVK DIEADKRRIV KRIPLHLTDT SKQTIVKSLA TLTEQLTVLF EEVQYDLVLI LGDRYEMLPV ANAALLYNIP ICHIHGGEKT MGNFDESIRH AITKMSHLHL TSTDEFRNRV IQLGENPTMY

Fig. 3 cont.

CPS2Q

SEQ. ID. NO. 25

charge margins

WO 00/05378

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PCT/NL99/00460

### 28/59

MELGIDFAED YYVVLFHPVT LEDNTAEEQT QALLDALKED GSQCLIIGSN SDTHADKIME LMHEFVKQDS DSYIFTSLPT RYYHSLVKHS QGLIGNSSSG LIEVPSLQVP TLNIGNRQFG RLSGPSVVHV GTSKEAIVGG LGQLRDVIDF TNPFEQPDSA LQGYRAIKEF LSVQASTMKE FYDR

Fig. 3 cont.

CPS2R

SEQ. ID. NO. 26

Change margino

PCT/NL99/00460

29/59

MKKVAFLGAG TFSDGVLPWL DRTRYELIGY FEDKPISDYR GYPVFGPLQD VLTYLDDGKV DAVFVTIGDN VKRKEIFDLL AKDHYDALFN IISEQANIFS PDSIKGRGVF

IGFSSFVGAD SYVYDNCIIN TGAIVEHHTT VEAHCNITPG VTINGLCRIG ESTYIGSGST

VIQCIEIAPY TTLGAGTVVL KSLTESGTYV GVPARKIK

Fig. 3 cont.

CPS2S

SEQ. ID. NO. 27

Change margins

MEPICLIPAR SGSKGLPNKN MLFLDGVPMI FHTIRAAIES GCFKKENIYV STDSEVYKEI CETTGVQVLM RPADLATDFT TSFQLNEHFL QDFSDDQVFV LLQVTSPLRS

GKHVKEAMEL YGKGQADHVV SFTKVDKSPT LFSTLDENGF AKDIAGLGGS YRRQDEKTLY

YPNGAIYISS KQAYLADKTY FSEKTAAYVM TKEDSIDVDD HFDFTGVIGR

IYFDYQRREQ QNKPFYKREL KRLCEQRVHD SLVIGDSRLL ALLLDGFDNI SIGGMTASTA

LENGGLFLAT PIKKVLLSLG VNDLITDYPL HMIEDTIRQL MESLVSKAEQ

VFVTTIAYTL FRDSVSNEEI VQLNDVIVQS ASELGISVID LNEVVEKEAM LDYQYTNDGL

HFNQIGQERV NQLILTSLTR

Fig. 3 cont.

CPS2T

SEQ. ID. NO. 28

change margins

31/59 WO-00/05378 ATCGCCAAAC GAAATTGGCA TTATTTGATA TGATAGCAGT TGCAATTTCT GCAATCTTAA CAAGTCATAT ACCAAATGCT GATTTAAATC GTTCTGGAAT TTTTATCATA ATGATGGTTC ATTATTTGC ATTTTTATA TCTCGTATGC CAGTTGAATT TGAGTATAGA GGTAATCTGA TAGAGTTTGA AAAAACATTT AACTATAGTA TAATATTTGC AATTTTTCTT ACGCAGTAT CATTTTTGTT GGAGAATAAT TTCGCACTTT CAAGACGTGG TGCCGTGTAT TTCACATTAA TAAACTTCGT TTTGGTATAC CTATTTAACG TAATTATTAA GCAGTTTAAG GATAGCTTTC TATTTTCGAC AATCTATCAA AAAAAGACGA TTCTAATTAC AACGGCTGAA CGATGGGAAA ATATGCAAGT TTTATTTGAA TCACATAAAC AAATTCAAAA AAATCTTGTT GCATTGGTAG TTTTAGGTAC AGAAATAGAT AAAATTAATT TATCATTACC GCTCTATTAT TCTGTGGAAG AAGCTATAGA GTTTTCAACA AGGGAAGTGG TCGACCACGT CTTTATAAAT CTACCAAGTG AGTTTTTAGA CGTAAAGCAA TTCGTTTCAG ATTTTGAGTT GTTAGGTATT GATGTAAGCG TTGATATTAA TTCATTCGGT TTTACTGCGT TGAAAAACAA AAAAATCCAA CTGCTAGGTG ACCATAGCAT TGTAACTTTT TCCACAAATT TTTATAAGCC TAGTCATATC ATGATGAAAC GACTTTTGGA TATACTCGGA GCGGTAGTCG GGTTAATTAT TTGTGGTATA GTTTCTATTT TGTTAGTTCC AATTATTCGT AGAGATGGTG GACCGGCTAT TTTTGCTCAG AAACGAGTTG GACAGAATGG ACGCATATTT ACATTCTACA AGTTTCGATC GATGTATGTT GATGCTGAGG AGCGCAAAAA AGACTTGCTC AGCCAAAACC AGATGCAAGG GTGGGTATGT TTTAAAATGG GAAAAACGAT CCTAGAATTA CTCCAATTGG ACATTTCATA CGCAAAAACA AGTTTAGACG AGTTACCACA GTTTTATAAT GTTTTAATTG GCGATATGAG TCTAGTTGGT ACACGTCCAC CTACAGTTGA TGAATTTGAA AAATATACTC CTGGTCAAAA GAGACGATTG AGTTTTAAAC CAGGGATTAC AGGTCTCTGG CAGGTTAGTG GTCGTAGTAA TATCACAGAC TTCGACGACG TAGTTCGGTT GGACTTAGCA TACATTGATA ATTGGACTAT CTGGTCAGAT ATTAAAATTT CGGTTCTTCA GGGGGACATT TGACTCACTT GTATTTGTTA AAACCGTTTT GGAAGGAAGA AGAACGTTTT TGGGTAACAT TTGATAAAGA GGATGCAAGA AGTCTTTTGA AGAATGAAAA AATGTATCCA TGTTACTTTC CAACAAATCG CAATCTCATT AATTTAGTGA AAAATACTTT CTTAGCTTTC AAAATTTTAC GTGATGAGAA ACCAGATGTT ATTATTTCAT CTGGTGCGGC CGTTGCTGTC CCCTTCTTTT ACATCGGAAA ACTATTTGGA GCAAAGACGA TTTATATTGA AGTATTTGAT CGAGTTAATA AATCTACATT AACTGGAAAA CTAGTTTATC CCGTAACAGA TATTTTTATT GTTCAGTGGG AAGAAATGAA GAAGGTATAT CCTAAATCTA TTAACTTGGG GAGTATTTTT TAATGATTTT TGTAACAGTA GGAACTCATG AACAACAGTT TAATCGATTG ATAAAAGAGA TTGATTTATT GAAAAAAAT GGAAGTATAA CCGACGAAAT ATTTATTCAA ACAGGATATT CTGACTATAT TCCAGAATAT TGCAAGTATA AAAAATTTCT CAGTTACAAA GAAATGGAAC AATATATTAA CAAATCAGAA GTAGTTATTT GCCACGGAGG CCCCGCTACT TTTATGAATT CATTATCCAA AGGAAAAAA CAATTATTGT TTCCTAGACA AAAAAAGTAT GGTGAACATG TAAATGATCA TCAAGTAGAG TTTGTAAGAA GAATTTTACA AGATAATAAT ATTTTATTTA TAGAAAATAT AGATGATTTG TTTGAAAAAA TTATTGAAGT TTCTAAGCAA ACTAACTTTA CATCAAATAA TAATTTTTTT TGTGAAAGAT TAAAACAAAT AGTTGAAAAA TTTAATGAGG ATCAAGAAAA TGAATAATAA AAAAGATGCA TATTTGATAA TGGCTTATCA TAATTTTTCT CAGATTTTAC TGGAGAGGGA TACAGATATT ATCATCTTCT CTCAGGAGAA TGCACACCAT TAGTTCCTTC AGAATACCTG TATAATTATT TTAAATATTC TCAGGATTTA TATGTTGAAT TTACAAAAGA TGAGCAAAAA TATAAAGAAA ATAGGATATA TGAACGAGTT AAATGTTACA GATTATTTCC TAATATATCA GAAAAAACTA TTGATAATGT ACTGTTTAGA ATTTTATTAA GAATGTATCG AGCTTTTGAA TACTATTTAC AAAGATTGTT GTTTATTGAT AGAATAAAAA ACATGGTCTA AGAATAAGAT TTGGTTCTAA TTGGGTTTCG CTTCCACATG ATTTTGTGGC AATTCTTTTA TCAAATGAAA ACGAAACAGC TTATTTATTT AAGTAATCTA AATGTCCAGA TGAACTATTT ATACAGACAA TTATAGAAAA ATATGAATTT TCAAATAGAT TATCTAAATA TGGAAATTTA AGATATATAA AGTGGAAAAA ATCAACATCT TCTCCTATTG TCTTTACAGA TGATTCTATT GATGAATTGC TAAATGCAAG AAATTTAGGT TTTTTATTTG CTAGAAAGTT AAAAATAGAA TAAATTATTT AAATATGACC CGGAATATTT TATTTTTAAG TACTTCTGGT TGATTATTT TATTCCAGAG CAAAAGTATG TATTTTTATT AATTTTATG AATTTAATTT TATTTCATAT AAAATTTTTG AAAACTAAGC TAATATTAAA AAATGAAATT TTATTGTTTT TATTATGGTC TATATTATGT TTTGTTTCAG TAGTCACAAG TATGTTTGTT GAAATAAATT TTGAAAGATT ATTTGCAGAT TTTACTGCTC CCATAATTTG GATTATTGCA ATAATGTATT ATAATTTGTA TTCATTTATA AATATTGATT ATAAAAAATT AAAAAATAGT ATCTTTTTTA GTTTTTTTAGT TTTATTAGGT ATATCTGCAT TGTATATTAT TCAAAATGGG AAAGATATTG TATTTTTAGA CAGACACCTT ATAGGACTAG ACTATCTTAT AACAGGCGTC AAAACAAGGT TGGTTGGCTT TATGAACTAT CCTACGTTAA ATACCACTAC AATTATAGTT TCAATTCCGT TAATCTTTGC ACTTATAAAA AATAAAATGC AACAATTTTT TTTCTTGTGT CTTGCTTTTA

change margins

TACCGATCTA TTTAAGTGGA TCGAGAATTG GTAGTTTATC GCTAGCAATA TTAATTATAT GCTTGTTATG GAGATATATA GGTGGAAAAT TTGCTTGGAT AAAAAAGCTA ATAGTAATAT TTGTAATACT ACTTATTATT TTAAATACTG AATTGCTTTA CCATGAAATT TTGGCTGTTT ATAATTCTAG AGAATCAAGT AACGAAGCTA GATTTATTAT TTATCAAGGA AGTATTGATA AAGTATTAGA AAACAATATT TTATTTGGAT ATGGAATATC CGAATATTCA GTTACGGGAA CTTGGCTCGG AAGTCATTCA GGCTATATAT CATTTTTTTA TAAATCAGGA ATAGTTGGGT TGATTTTACT GATGTTTTCT TTTTTTTATG TTATAAAAAA ACATCATTAG CCATATTTTT CATATAGAA ACAATAGATC CGATTATTAT TATATTAGTA CTATTCTTTT CTTCAATAGG TATTTGGAAT AATATAAATT TTAAAAAGGA TATGGAGACA AAAAATGAAT GATTTAATTT CAGTTATTGT ACCAATTTAT AATGTCCAAG ATTATCTTGA TAAATGTATT AACAGTATTA TTAACCAAAC ATATACTAAT TTAGAGGTTA TTCTCGTAAA TGATGGAAGT ACTGATGATT CTGAGAAAAT TTGCTTAAAC TATATGAAGA ACGATGGAAG AATTAAATAT TACAAGAAAA TTAATGGCGG TCTAGCAGAT GCTCGAAATT TCGGACTAGA ACATGCAACA GGTAAATATA TTGCTTTTGT CGATTCTGAT GACTATATAG AAGTTGCAAT GTTCGAGAGA ATGCATGATA ATATAACTGA GTATAATGCC GATATAGCAG AGATAGATTT TTGTTTAGTA GACGAAAACG GGTATACAAA GAAAAAAAGA AATAGTAATT TTCATGTCTT AACGAGAGAA GAGACTGTAA AAGAATTTTT GTCAGGATCT AATATAGAAA ATAATGTTTG GTGCAAGCTT TATTCACGAG ATATTATAAA AGATATAAAA TTCCAAATTA ATAATAGAAG TATTGGTGAG GATTTGCTTT TTAATTTGGA GGTCTTGAAC AATGTAACAC GTGTAGTAGT TGATACTAGA GAATATTATT ATAATTATGT CATTCGTAAC AGTTCGCTTA TTAATCAGAA ATTCTCTATA AATAATATTG ATTTAGTCAC AAGATTGGAG AATTACCCCT TTAAGTTAAA AAGAGAGTTT AGTCATTATT TTGATGCAAA AGTTATTAAA GAGAAGGTTA AATGTTTAAA CAAAATGTAT TCAACAGATT GTTTGGATAA TGAGTTCTTG CCAATATTAG AGTCTTATCG AAAAGAAATA CGTAGATATC CATTTATTAA AGCGAAAAGA TATTTATCAA GAAAGCATTT AGTTACGTTG TATTTGATGA AATTTTCGCC TAAACTATAT GTAATGTTAT ATAAGAAATT TCAAAAGCAG TAGAGGTAAA AATGGATAAA ATTAGTGTTA TTGTTCCAGT TTATAATGTA GATAAATATT TAAGTAGTTG TATAGAAAGC ATTATTAATC AAAATTATAA AAATATAGAA ATATTATTGA TAGATGATGG CTCTGTAGAT GATTCTGCTA AAATATGCAA GGAATATGCA GAAAAAGATA AAAGAGTAAA AATTTTTTC ACTAATCATA GTGGAGTATC AAATGCTAGA AATCATGGAA TAAAGCGGAG TACAGCTGAA TATATTATGT TTGTTGACTC TGATGATGTT GTTGATAGTA GATTAGTAGA AAAATTATAT TTTAATATTA TAAAAAGTAG AAGTGATTTA TCTGGTTGTT TGTACGCTAC TTTTTCAGAA AATATAAATA ATTTTGAAGT GAATAATCCA AATATTGATT TTGAAGCAAT TAATACCGTG CAGGACATGG GAGAAAAAA TTTTATGAAT TTGTATATAA ATAATATTTT TTCTACTCCT GTTTGTAAAC TATATAAGAA AAGATACATA ACAGATCTTT TTCAAGAGAA TCAATGGTTA GGAGAAGATT TACTTTTTAA TCTGCATTAT TTAAAGAATA TAGATAGAGT TAGTTATTTG ACTGAACATC TTTATTTTTA TAGGAGAGGT ATACTAAGTA CAGTAAATTC TTTTAAAGAA GGTGTGTTTT TGCAATTGGA AAATTTGCAA AAACAAGTGA TAGTATTGTT TAAGCAAATA TATGGTGAGG ATTTTGACGT ATCAATTGTT AAAGATACTA TACGTTGGCA AGTATTTTAT TATAGCTTAC TAATGTTTAA ATACGGAAAA CAGTCTATTT TTGACAAATT TTTAATTTTT AGAAATCTTT ATAAAAAATA TTATTTTAAC TTGTTAAAAG TATCTAACAA AAATTCTTTG TCTAAAAATT TTTGTATAAG AATTGTTTCG AACAAAGTTT TTAAAAAAT ATTATGGTTA TAATAGGAAG ATATCATGGA TACTATTAGT AAAATTTCTA TAATTGTACC TATATATAAT GTAGAAAAAT ATTTATCTAA ATGTATAGAT AGCATTGTAA ATCAGACCTA CAAACATATA GAGATTCTTC TGGTGAATGA CGGTAGTACG GATAATTCGG AAGAAATTTG TTTAGCATAT GCGAAGAAAG ATAGTCGCAT TCGTTATTTT AAAAAAGAGA ACGGCGGGCT ATCAGATGCC CGTAATTATG GCATAAGTCG CGCCAAGGGT GACTACTTAG CTTTTATAGA CTCAGATGAT TTTATTCATT CGGAGTTCAT CCAACGTTTA CACGAAGCAA TTGAGAGAGA GAATGCCCTT GTGGCAGTTG CTGGTTATGA TAGGGTAGAT GCTTCGGGGC ATTTCTTAAC AGCAGAGCCG CTTCCTACAA ATCAGGCTGT TCTGAGCGGC AGGAATGTTT GTAAAAAGCT GCTAGAGGCG GATGGTCATC GCTTTGTGGT GGCCTGTAAT AAACTCTATA AAAAAGAACT ATTTGAAGAT TTTCGATTTG AAAAGGGTAA GATTCATGAA GATGAATACT TCACTTATCG CTTGCTCTAT GAGTTAGAAA AAGTTGCAAT AGTTAAGGAG TGCTTGTACT ATTATGTTGA CCGAGAAAAT AGTATCACAA CTTCTAGCAT GACTGACCAT CGCTTCCATT GCCTACTGGA ATTTCAAAAT GAACGAATGG ACTTCTATGA AAGTAGAGGA GATAAAGAGC TCTTACTAGA GTGTTATCGT TCATTTTTAG CCTTTGCTGT TTTGTTTTTA GGCAAATATA ATCATTGGTT GAGCAAACAG CAAAAGAAGC TT

Fig. 4 cont.

modify inaguno

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33/59

| ROTKLALFDM       | IAVAISAILT        | SHIPNADLNR | SGIFIIMMVH | YFAFFISRMP | VEFEYRGNLI |
|------------------|-------------------|------------|------------|------------|------------|
| EFEKTFNYSI       | <b>IFAIFLTAVS</b> | FLLENNFALS | RRGAVYFTLI | NFVLVYLFNV |            |
| IIKOFKDSFL       | FSTIYQKKTI        | LITTAERWEN | MQVLFESHKQ | IQKNLVALVV | LGTEIDKINL |
| SLPLYYSVEE       | <b>AIEFSTREVV</b> | DHVFINLPSE | FLDVKQFVSD | FELLGIDVSV |            |
| DINSFGFTAL       | KNKKIQLLGD        | HSIVTFSTNF | YKPSHIMMKR | LLDILGAVVG | LIICGIVSIL |
| LVPIIRRDGG       | PAIFAQKRVG        | QNGRIFTFYK | FRSMYVDAEE | RKKDLLSQNQ |            |
| MOGWVCFKMG       | KTILELLQLD        | ISYAKTSLDE | LPQFYNVLIG | DMSLVGTRPP | TVDEFEKYTP |
| GOKRRLSFKP       | GITGLWQVSG        | RSNITDFDDV | VRLDLAYIDN | WTIWSDIKIL |            |
| T PARTACE TO THE | FCSK              |            |            |            |            |

Fig. 4 cont.

CPS1E

SEQ. ID. NO. 30

Modify margins

MKVCLVGSSG GHLTHLYLLK PFWKEEERFW VTFDKEDARS LLKNEKMYPC YFPTNRNLIN LVKNTFLAFK ILRDEKPDVI ISSGAAVAVP FFYIGKLFGA KTIYIEVFDR VNKSTLTGKL VYPVTDIFIV QWEEMKKVYP KSINLGSIF

Fig. 4 cont.

CPS1F

SEQ. ID. NO. 31

midity margino

WO 00/05378

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PCT/NL99/00460

MIFVTVGTHE QQFNRLIKEI DLLKKNGSIT DEIFIQTGYS DYIPEYCKYK KFLSYKEMEQ YINKSEVVIC HGGPATFMNS LSKGKKQLLF PRQKKYGEHV NDHQVEFVRR ILQDNNILFI ENIDDLFEKI IEVSKQTNFT SNNNFFCERL KQIVEKFNED QENE

Fig. 4 cont.

CPS1G

SEQ. ID. NO. 32

molity margins

|            |            |            | ′59 J      |            |            |
|------------|------------|------------|------------|------------|------------|
| MFKLFKYDPE | YFIFKYFWLI | IFIPEQKYVF | LLIFMNLILF | HIKFLKTKLI | LKNEILLFLL |
| WSILCFVSVV | TSMFVEINFE | RLFADFTAPI | IWILAIMYYN | LYSFINIDYK |            |
| KLKNSIFFSF | LVLLGISALY | IIQNGKDIVF | LDRHLIGLDY | LITGVKTRLV | GFMNYPTLNT |
| TTIIVSIPLI | FALIKNKMQQ | FFFLCLAFIP | IYLSGSRIGS | LSPLAILIIC |            |
| LLWRYIGGKF | AWIKKLIVIF | VILLIILNTE | LLYHEILAVY | NSRESSNEAR | FIIYQGSIDK |
| VLENNILFGY | GISEYSVTGT | WLGSHSGYIS | FFYKSGIVGL | ILLMFSFFYV | _          |
| IKKSYGVNGE | TALFYFTSLA | IFFIYETIDP | IIIILVLFFS | SIGIWNNINF | KKDMETKNE  |

Fig. 4 cont.

CPS1H

SEQ. ID. NO. 33

modefy margins

| _ | /59 |   |
|---|-----|---|
|   | /54 | ė |

|             |            | 37,        | /59 4      |            |            |
|-------------|------------|------------|------------|------------|------------|
| MNDI.TSVIVP | IYNVODYLDK | CINSIINQTY | TNLEVILVND | GSTDDSEKIC | LNYMKNDGRI |
| KYYKKTNGGL  | ADARNEGLEH | ATGKYIAFVD | SDDYIEVAMF | ERMHDNITEY |            |
| NADIAEIDFC  | LVDENGYTKK | KRNSNFHVLT | REETVKEFLS | GSNIENNVWC | KLYSRDIIKD |
| IKFQINNRSI  | GEDLLFNLEV | LNNVTRVVVD | TREYYYNYVI | RNSSLINQKF |            |
| SINNIDLVTR  | LENYPFKLKR | EFSHYFDAKV | IKEKVKCLNK | MYSTDCLDNE | FLPILESYRK |
| EIRRYPFIKA  | KRYLSRKHLV | TLYLMKFSPK | LYVMLYKKFQ | KQ         |            |

Fig. 4 cont.

CPS1I

SEQ. ID. NO. 34

modely margins

| 38 | /59  | J        | , |
|----|------|----------|---|
| 30 | / 22 | $\sim 1$ | 1 |

|            |            |            | \ <i>U</i> |            |            |
|------------|------------|------------|------------|------------|------------|
| MDKISVIVPV | YNVDKYLSSC | IESIINQNYK | NIEILLIDDG | SVDDSAKICK | EYEKDKRVKI |
| FFTNHSGVSN | ARNHGIKRST | AEYIMFVDSD | DVVDSRLVEK | LYFNIIKSRS |            |
| DLSGCLYATF | SENINNFEVN | NPNIDFEAIN | TVQDMGEKNF | MNLXXNNIFS | TPVCXLYQKR |
| YITDLFQENQ | WLGEDLLFNL | HYLKNIDRVS | YLTEHLYFYR | RGILSTVNSF |            |
| KEGVFLQLEN | LQKQVIVLFK | QIYGEDFDVS | IVKDTIRWQV | FYYSLLMFKY | GKQSIFDKFL |
| IFRNLYKKYY | FNLLKVSNKN | SLSKNFCIRI | VSNKVFKKIL | WL         |            |

Fig. 4 cont.

CPS1J

SEQ. ID. NO. 35

modify marguns

PCT/NL99/00460

39/59

MDTISKISII VPIYNVEKYL SKCIDSIVNO TYKHIEILLV NDGSTDNSEE ICLAYAKKOS

RIRYFKKENG GLSDARNYGI SRAKGDYLAF IDSDDFIHSE FIQRLHEAIE

RENALVAVAG YDRVDASGHF LTAEPLPTNQ AVLSGRNVCK KLLEADGHRF VVACNKLYKK

ELFEDFRFEK GKIHEDEYFT YRLLYELEKV AIVKECLYYY VDRENSITTS

SMTDHRFHCL LEFQNERMDF YESRGDKELL LECYRSFLAF AVLFLGKYNH WLSKQQKK

Fig. 4 cont.

CPS1K

SEQ. ID. NO. 36

modify manzins

## 40/59

|                          |                          | 40)         |                               |                          |                    |
|--------------------------|--------------------------|-------------|-------------------------------|--------------------------|--------------------|
| AAGCTTATCG               |                          |             |                               | TCATAGACGA               | AAAGGGATGT         |
|                          | AGAAAAAGTT               |             | ACTTTCTTCA                    |                          | C                  |
|                          | AAGTTTATCC               | TGAAATACGA  |                               | GTGCTGAATT               | GTATTATAGT         |
| AAAGATATAT               |                          |             | AAAGTACCCA                    |                          | MMCN NCN NCC       |
|                          | ATTCTTTTGG               |             |                               | TGGAAAGAGA               | TTCAAGAAGC         |
|                          | GTGACGCTAC               |             |                               | GCCCATATAG               | GACAAGGGAT         |
|                          | CGCCCTAGCG               |             |                               | AGAGTTAATT               | GACAAGGGAI         |
|                          | GGTAAATAGT               |             |                               | TTTAATTGGT               | TTTAGTACAT         |
|                          | AAGAATTTAA<br>GCGATATGCA |             | CGGTATTTT                     | CGTTTATGAG               | IIIAGIACAI         |
|                          |                          | CAGAGGAATT  | AGTAGACCTC                    | AAAGCGAAAG               | CGTTGCTAAA         |
| GGAGGCTTAT<br>AAAGAATCCT |                          |             | GGCGATTTAA                    |                          | COTTOCTIBET        |
|                          | GAGAGAAAA                |             |                               | GATAAACTGT               | TAGAACGCAA         |
| CAGTAAACGA               |                          |             |                               | CTTATAGTTT               | 11101111000111     |
| CCATGATTT                |                          |             |                               | CATACCAGAT               | GAACGCTTCA         |
| TTCTTGCAGT               |                          |             |                               | ATCGTTTAGA               |                    |
| TTAAAAGTCT               |                          |             |                               | AGAGTTATGT               | AAAAATAGGA         |
|                          | TATCTGCGCA               |             |                               | CAATGGTGTT               |                    |
| GTGGCAGGCT               |                          |             | AGTATCCTTA                    |                          | ATGTAATGCT         |
|                          | AGGATTGTTT               |             |                               | AGAAAAAATG               |                    |
|                          | GAAGGATAGC               |             |                               | AGGTGCTGGA               | GATGGTGGTA         |
| ATATTTTTAT               |                          | AAAGATCGAA  | AATTGAATTT                    | TGAAATTGTC               |                    |
| GGTATCGTTG               | ATCGTGATCC               | AAATAAACTT  |                               | TCCGTACGGC               | TAAAGTTTTA         |
| GGAAACCGTA               | ATGATATTCC               | ACGACTGGTA  | GAGGAATTAG                    | CTGTTGACCA               |                    |
| AGTGACGATT               | GCCATCCCTT               | CTTTAAATGG  |                               | GAGAAGATTG               | TTGAAATCTG         |
|                          | GGAGTGACCG               |             |                               | GAAGACATTA               |                    |
| TGGCGGGGAA               | CATGTCTGTC               | AGTGCCTTTC  |                               | CGTAGCAGAC               | CTTCTTGGTC         |
| GACCAGAGGT               | TGTTTTGGAT               |             | TGAATCAGTT                    | TTTCCAAGGG               |                    |
| AAAACAATCC               |                          | AGCAGGTGGC  | TCTATCGGTT                    | CAGAGCTATG               | TCGTCAAATT         |
| GCTAAGTTTA               | CGCCTAAACG               | CTTGTTGTTG  | CTTGGACATG                    | GAGAAAATTC               | N COMMCCONCCC      |
|                          | ATTCATCGAG               |             | AAAGTACCAA                    | GGTAAGATTG               | AGTTGGTCCC         |
|                          | GATATTCAAG               |             |                               | ATAATGGCTG               | TTGATGGAAT         |
| AATATCAACC               | CGATGTTGTT               | TATCATGCTG  |                               | GCATGTTCCT<br>GAAGAATGTG | IIGAIGGAAI         |
| ATAATCCACA               | TGAAGCAGTG               | AAGAAIAAIA  |                               | TGGTTTCAAC               | AGATAAAGCT         |
| GCTGAGGCGG               | CAAATGTCAT               | CCCACCCACT  | AAATTIGITA                    | CAGAAATGAT               |                    |
| TGTTACAGGT               |                          | CAGGTCAGAC  | TCAATTTGCG                    | GCAGTCCGGT               | TTGGGAATGT         |
| TCTAGGTAGT               | CGTGGAAGTG               |             | ATTCAAAGAG                    |                          |                    |
| AAGGTGGACC               |                          | ACCGACTTTA  |                               |                          | ACGATTCCTG         |
| AGGCAAGTCG               |                          | CAAGCTGGAC  | ATTTGGCAAA                    |                          |                    |
| ATATTTGTCT               | TGGATATGGG               |             |                               | AATTGGCAAG               | AAAAGTTATC         |
| TTGTTAAGTG               | GACACACAGA               | GGAAGAAATC  | GGGATTGTAG                    | AATCTGGAAT               |                    |
| CAGACCAGGC               | GAGAAACTCT               | ACGAGGAATT  | ATTATCAACA                    | GAAGAACGTG               | TCAGCGAACA         |
| CATTCATGAA               | AAAATATTTG               | TGGGTCGCGT  | TACAAATAAG                    | CAGTCGGACA               | •                  |
| TTGTCAATTC               | ATTTATCAAT               | GGATTACTCC  | AAAAAGATAG                    | AAATGAATTA               | AAAAATATGT         |
| TGATTGAATT               | TGCAAAACAA               | GAATAAGAAA  | GTAAAAAATA                    | TTTTTACTTT               |                    |
| CCTAGAGTTT               | AAACGATGTT               | TAAGTTCTAG  | GAAGGTTAGA                    | ATACCTAATT               | AACAACAATA         |
| TTACTATTTA               | TTAAGAGTCA               | GATAATAGCA  | ACTAAGTGCT                    | ACAAACTATC               |                    |
| TTTATAATAA               | GTATATTTGG               | TCAAAAGGGA  | GATGTGAAAT                    | GTATCCAATT               | TGTAAACGTA         |
| TTTTAGCAAT               | TATTATCTCA               | GGGATTGCTA  | TTGTTGTTCT                    | GAGTCCAATT               |                    |
| TTATTATTGA               | TTGCATTGGC               | AATTAAATTA  | GATTCTAAAG                    | GTCCGGTATT               | ATTTAAACAA         |
| AAGCGGGTTG               | GTAAAAACAA               | GTCATACTTT  | ATGATTTATA                    | AATTCCGTTC               | CHARCCCCAT         |
| TATGTACGTT               | GACGCACCAA               | GTGATATGCC  | GACTCATCTA                    | TTAAAGGATC               | CTAAGGCGAT         |
| GATTACCAAG               | GTGGGCGCGT               | TTCTCAGAAA  | AACAAGTTTA                    | GATGAACTGC               | <b>ででかてでで町町が</b> 町 |
| CACAGCTTTT               | TAATATTTTT               | AAAGGTGAAA  | TGGCGATTGT                    | TGGTCCACGC               | CCAGCCITAI         |
| GGAATCAATA               | TGACTTAATT               | GAAGAGCGAG  | ATAAATATGG                    | CTCATAIGAI               | ССАВАТТСВТ         |
| ATTCGTCCTG               | GACTAACCGG               | TTGGGCTCAA  | ATTAATGGTC                    | GTGATGAATT               | GGMMIIGHI          |
| GAAAAGTCAA               | AATTAGATGG               | ATATTATGTT  | CAMMAIAIGA                    | DCDDGCCDDG               | СФСФФСФФСА         |
| GGATATTAAA               | GGGCAGAAAG               | CANANCCAMC  | CHGIGIHGCC<br>A A A TOTTO THE | AGAAGCGAAG               | CIGITOTION         |
| AGGTGGAACA               | DACARAAAG                | CACACMMMCM  | TACCCA A TOTAL                | TTGGAAAGCA               | ТССТТСТСАА         |
| GGTCTATGAG               | ATTCCAACGG               | ACCHIOCHCIT | TAGGGWATCI                    | GGGCCACTCA               | 1001101011         |
| TCAAACAATG               | ATTUCAAUGG               | TOUTIGICIT  | TTADADACTCC                   | ATTTTCATTT               | TTTAAAACGA         |
| ATCAGAGCTT               | AAAGAATTCG               | TINGHMORAL  | TIMMMGICG                     | TGAAGGTTTG               |                    |
| TAGCCTTGGA               | AMMUMMIIUG               | CCTTTAGGAA  | AAATGCACIGAA                  | TGATGATGTT               | GCATATACAT         |
| AAACATTGTA               | AAAAGCAAGT               | TAACTTACACG | A A A C A A A A A C C         | CCACTATAGA               |                    |
| ACACGTTTTG               | NANAUCAAU I              | A           | A -                           | seletris mi              | 1100 MO            |
|                          |                          | 1           | (2.2.7                        | secur us lile            | William Co.        |

|       | 1   |
|-------|-----|
| 1/59  | J   |
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|            |            |            | -          |                   | , , ,      |
|------------|------------|------------|------------|-------------------|------------|
| TATTGAGATA | GATGAGTTCT | TAAATTCTAC | TAGTGAAATA | GTTTCTCATA        | AAAATGTTCC |
| AACCCAGCAC | GATGAAATAT | TAAAGATGGC | AAGGCGGGAG | AAATCCATGT        |            |
| GCCACATGAC | TGTAATGTTT | AAAAAGAAAA | GTGTCGAGAG | AGCAGGGGGG        | TATCAAACAC |
| TTCCGTACGT | AGAAGATTAT | TTCCTTTGGG | TGCGCATGAT | TGCTTCAGGA        |            |
| TCGAAATTTG | CAAACATTGA | TGAAACACTA | GTTCTTGCAC | GTGTTGGAAA        | TGGGATGTTC |
| AATAGGAGGG | GGAACAGAGA | ACAAATTAAC | AGTTGGACAT | TACTAATTGA        |            |
| ATTTATGTTA | GCTCAAGGAA | TTGTTACACC | ACTAGATGTA | TTTATTAATC        | AAATTTACAT |
| TAGGGTCTTT |            | CAACTTGGAT |            | ATTTATGGAA        |            |
| AAATCTTAAG | GAAATAGTAT | GATTACAGTA | TTGATGGCTA | CATATAATGG        | AAGCCCATTT |
| ATAATAAAAC | AGTTAGATTC | AATTCGAAAT | CAAAGTGTAT | CAGCAGACAA        |            |
| AGTTATTATT | TGGGATGATT | GCTCGACAGA | TGATACAATA | AAAATAATAA        | AAGATTATAT |
| AAAAAAATAT | TCTTTGGATT | CATGGGTTGT | CTCTCAAAAT | AAATCTAATC        |            |
| AGGGGCATTA | TCAAACATTT | ATAAATTTGA | CAAAGTTAGT | TCAGGAAGGA        | ATAGTCTTTT |
| TTTCAGATCA |            | TGGGACTGTC | ATAAAATTGA | GACAATGCTT        |            |
| CCAATCTTTG | ACAGAGAAAA | TGTATCAATG | GTGTTTTGCA | <b>AATCCAGATT</b> | GATTGATGAA |
|            | TTATCAGTAG | CCCAGATACT | TCGGATAGAA | TCAATACGTA        |            |
| CTCTCTAGA  |            |            |            |                   |            |
| 0.0.01110  |            |            |            |                   |            |

Fig. 5 cont.

SEQ. ID. NO. 37

modify margins

PCT/NL99/00460

42/59

12/39

AYROGVRYIV ATSHRRKGMF ETPEKVIMTN FLOFKDAVAE VYPEIRLCYG AELYYSKDIL

SKLEKKKVPT LNGSRYILLE FSSDTPWKEI QEAVNEVTLL GLTPVLAHIE

RYDALAFHAE RVEELIDKGC YTQVNSNHVL KPTLIGDRAK EFKKRTRYFL EQDLVHCVAS

DMHNLSSRPP FMREAYKLLT EEFGKDKAKA LLKKNPLMLL KNQAI

Fig. 5 cont.

CPS9D

SEQ. ID. NO. 38

modefy margins

|                      |                      |             |            | 1           | · · · · · · · · · · · · · · · · · · · |
|----------------------|----------------------|-------------|------------|-------------|---------------------------------------|
|                      |                      | 43,         | /59        | 6           | •                                     |
| MOT CONTONIA.        | LERNSKRLIL           | VCMDTCLLIV  | SMILSRLFLD | VIIDIPDERF  | ILAVLFVSIL                            |
| TO THE OWNER AND THE | CCT TTDVTCY          | OSVVKTGLST. | ISAHSLFLII | SMVLWQArSi  |                                       |
| DETT VELETS          | YVMI.TTPRIV          | WKVLHETRKN  | AIRKKDSPLR | TTA AGMGDGG | NIFINTVKDR                            |
|                      | DEDDNKTCTE           | TRTAKVI.GNR | NDTPRLVEEL | AADOALTHIE  |                                       |
| OT NOVEDEKT          | VETCNTTGVT           | VNNMPSIEDI  | MAGNMSVSAF | OFIDAMPITE  | RPEVVLDQDE                            |
| - NOTE OCIONE        | TUTCACCSTC           | SELCROTAKE  | TPKRLLLLGH | GENZITTIK   |                                       |
| DI I EVVOCKI         | FI.VPI.TADTO         | DRELIFSIMA  | EYOPDVVYHA | WAHVUALTE   | YNPHEAVKNN                            |
|                      | <b>カシͲカレス/カレビス</b> / | MUCTUKTUND  | PNVMGATKRV | AEMIALGINE  |                                       |
| DOOMOEN NVD          | FCNVLGSRGS           | VVPLFKEOIR  | KGGPVTVTDE | KMIKILMITE  | FYSKTATÖVQ                            |
| ··· PROCETES         | T DMCFDVATI.         | FIARKVILLS  | GHTEEEIGIV | FOCIKEGEVT  |                                       |
| YEELLSTEER           | VSEQIHEKIF           | VGRVTNKQSD  | IVNSFINGLL | QKDRNELKNM  | LIEFAKOE                              |

Fig. 5 cont.

CPS9E

SEQ. ID. NO. 39

modify margino

WO 00/05378

MYPICKRILA IIISGIAIVV LSPILLIAL AIKLDSKGPV LFKQKRVGKN KSYFMIYKFR

SMYVDAPSDM PTHLLKDPKA MITKVGAFLR KTSLDELPQL FNIFKGEMAI

VGPRPALWNQ YDLIEERDKY GANDIRPGLT GWAQINGRDE LEIDEKSKLD GYYVQNMSLG

LDIKCFLGTF LSVARSEGVV EGGTGQKGKG

Fig. 5 cont.

CPS9F

SEQ. ID. NO. 40

Modefy margins 45/59

1

|            | 43/39      |            |            |           |            |  |  |
|------------|------------|------------|------------|-----------|------------|--|--|
| MKFSVLMSVY |            |            |            |           | ILEEFKSRFS |  |  |
| FFKTIALEKN |            |            |            |           |            |  |  |
|            |            |            |            |           | FKKKSVERAG |  |  |
| GYQTLPYVED |            |            |            |           |            |  |  |
| NSWTLLIEFM | LAOGIVTPLD | VFINOIYIRV | FVYMPTWIKK | LIYGKILRK |            |  |  |

Fig. 5 cont.

CPS9G

SEQ. ID. NO. 41

modify margins

46/59
MITVLMATYN GSPFIIKQLD SIRNQSVSAD KVIIWDDCST DDTIKIIKDY IKKYSLDSWV :

VSONKSNOGH YOTFINLTKL VQEGIVFFSD QDDIWDCHKI ETMLPIFDRE

NVSMVFCKSR LIDENGNIIS SPDTSDRINT YSL

Fig. 5 cont.

CPS9H

SEQ. ID. NO. 42

modefy margins

| W <del>O 00/05378</del>   |                             | 47/            | 59                     | $\sim$                        | PCT/NL99/00460      |
|---|-----------------------------|----------------|------------------------|-------------------------------|---------------------|
| CTGCAGCACA  | maaccamcmm /                | ССВФФСВФСС     | дататаатсс             | ACATGAAGCA                    |                     |
| CTGCAGCACA  | AACGAAGAAT                  | CORTIGRIGG .   | CCCCTAAAAC             | TCCAAAGGTT                    | ů.                  |
| ATATTTTTGG  | AACGAAGAAT                  | GIGGCIGAGG     | CGGCIAAAAC             | CCCCAAATGT                    | CATGGGAGCG          |
| GCCAAATTTG  | TTATGGTTTC                  | AACAGAIAAA     | GCIGIIMAIC             | ACCCAGGTCA                    |                     |
| ACTAAACGTG  | TTGCAGAAAT                  | GATTGTAACA     | GGTTTAAACG             | ACCCAGGICA                    | СТСТТСТТСС          |
| GACTCAATTT  | GCGGCAGTCC                  | GTTTTGGGAA     | TGTTCTAGGT             | AGTCGTGGAA                    | GIGIIGIICO          |
| GCTATTCAAA  | GAGCAAATTA                  | GAAAAGGTGG     | ACCTGTTACG             | GTTACCGACT                    | A MCCA A CCMC       |
| TTAGGATGAC  | TCGTTATTTC                  | ATGACGATTC     | CTGAGGCAAG             | TCGTTTGGTT                    | ATCCAAGCTG          |
| これのみの中でのこと  | AAAAGGTGGA                  | GAAATCTTTG     | TCTTGGATAT             | GGGTGAGCCA                    |                     |
| ርጥክር ል ል ልጥር ር  | TGGAATTGGC                  | AAGAAAAGTT     | ATCTTGTTAA             | GCGGACATAC                    | AGAGGAAGAA          |
| カのこことの大型では  | <b>ТАСААТСТСС</b>           | AATCAGACCA     | GGCGAGAAAC             | TCTACGAGGA                    |                     |
| አጥጥር:ሞጥልጥCA   | ACAGAAGAAC                  | GTGTCAGCGA     | ACAGATTCAT             | GAAAAAATAT                    | TTGTGGGTCG          |
|   | NACCACTCCC                  | ACATTGTCAA     | TTCATTTATC             | AATGGATTAC                    |                     |
| σοσανάδο  | TAGAAATGAA                  | TTAAAAGATA     | TGTTGATTGA             | ATTTGCAAAA                    | CAAGAATAAG          |
| 777CM77777  | <b>አ</b> ጥከጥጥጥጥልር           | TTTCCTAGAG     | TTTAAACGAT             | GTTTAAGTTC                    |                     |
| MANGIAMM  | GGAATTGCTT                  | TCGTGGAGGT     | GATAGATAGA             | <b>AACCTATATA</b>             | TTTGTAGAAG          |
| TAGGAAGGII  | AAACTAAAGG                  | TCDATCGGAA     | CATAAAGTTT             | AGATAGAGTT                    |                     |
| AAAGGATATT  | CCCNNACACC                  | TCAATCCAAC     | CTCTCGCTCG             | TTACTAAGCA                    | GGAGATAGTA          |
| GGTATTTAAT  | AAAGAGAGTT                  | TGAMIGCAAC     | TATAACTAGG             | CTAAAGTGAG                    |                     |
| AAGTTGCTTG  | AAAGAGAGTT                  | 1G11MA1CAG     | CTATATOTACA            | CAATTATTGT                    | AGTGGGGATA          |
| AATATATATC  | TATTATTATC                  | GGTAATGATA     | CTATIATIOA             | CCTTABABAA                    |                     |
| AAAATAATTT  | TTGGTGATTT                  | TATCGTCCGA     | THAMAGGIG              | <b>ጥጥልጥልልጥልጥ</b> ጥ            | TATAGGAGAT          |
| GTACTTATAT  | TCTTTTAGAA                  | TTGATGAAAA     | ATAIGGGGA              | ACARCABTCA                    | TATAGGAGAT          |
| ACGATGACTA  | GAGTAGAGTT                  | GATTACTAGA     | GAATTTTTA              | AGMAGMATGA<br>CCTC N N TT N T | ጥጥልጥጥልልልጥጥ          |
| AGCAACCAGT  | AAATATTTTC                  | AGAAGA'I'AGA   | ATCAAGAAGA             | GGIGAAIIAI                    | TTATTAAATT          |
| CTTTATGGAT  | AAGTTACTTG                  | CGCTTATCCT     | ATTATTGCTA             | TTATCCCCAG                    | መመመጥ <b>አ</b> ጥርርርር |
| TAATCATTAT  | ATTAGCTATT                  | TGGATAAAAT     | TAGATAGTAA             | GGGGCCAAII                    | TTTTATCGCC          |
| * * C * * C C T C T   | <b>ጥአ</b> ሮር <b>አር</b> ልጥልጥ | CCTCCAATTT     | TTAGAATATT             | TAAGIIIAGA                    |                     |
| ACAATGATTT  | CTGATGCGGA                  | TAAAGTCGGA     | AGTCTTGTCA             | CAGTCGGTCA                    | AGATAATCGT          |
| *************   | <b>ጥ</b> ሮርርጥሮ እር እጥ        | ጥልጥሮልርልልልል     | TATCGGCTGG             | ACGAAGIGCC                    |                     |
| <b>ሮሮ</b> እስርጥጥጥጥ   | ΔΔΤΩΤΤΤΤΑΑ                  | TGGGGGATAT     | GAGCTTTGTA             | GGTGTAAGAC                    | CAGAAGTACA          |
| ካ ካ ካ ካ ጠ ካ መ <i>ር</i> ጥ እ  | <b>አአጥሮ</b> ልሮሞ <b>ል</b> ሞል | CTGATGAAAT     | GTTTGCGACG             | TTACTTTTAC                    |                     |
| CTCCACCAAT  | TACTTCACCA                  | GCGAGTATTG     | CATATAAGGA             | TGAAGATATI                    | GIIIIAGAAG          |
| $\mathbf{x}$ $\mathbf{x}$ $\mathbf{m}$ $\mathbf{x}$ $\mathbf{m}$ $\mathbf{m}$ $\mathbf{c}$ $\mathbf{m}$ $\mathbf{m}$ $\mathbf{c}$ | TCDACCCTAT                  | AGTCCTGATG     | AAGCATATGT             | TCAAAAAGTA                    | •                   |
| ͲͲΝϹϹΝGΔΔΑ  | AAATGAAGTA                  | CAATTTGGAA     | TATATCAGAA             | ACTITGGAAT                    | TATTICIGAL          |
| MMM 7 7 CM7 7   | ጥሮአጥጥርልጥልሮ                  | ΔατάδτταλΑ     | GTAATAAAAT             | AGGAGATTAA                    | •                   |
| <b>ΑΛΨΩΛΓΑΔΑΑ</b>   | AGACAAAATA                  | TTCCATTTTC     | ACCACCAGAT             | ATTACCCAAG                    | CTGAAATIGA          |
| መር እ እ ርመመ እጥጥ  | CACACACTAA                  | AATCTGGTTG     | GATTACAACA             | GGACCAAAGA                    | L                   |
| CAAAACACCT  | AGAACGTCGG                  | CTATCAGTAT     | TTACAGGAAC             | CAATAAAACI                    | GIGIGIIIAA          |
| N MM CMCCMNC  | TOCAGGATTG                  | CAACTAGTCT     | TACGAATTCT             | TGGTGTTGGA                    |                     |
| ATTOTOGIAC  | DACTTATTCT                  | TCCTGCTATG     | ACCTATACTG             | CCTCATGTAG                    | TGTCATTACT          |
| CABCEACCAC  | CAACTCCTGT                  | GATGGTTGAT     | ATTCAAAAAA             | ACAGCTTTGA                    |                     |
| CATGIAGGAG  | CAMCICCIOICI                | AAAAAGCGAT     | TACTCCGAAA             | ACAAAAGTTA                    | TCATTCCTGT          |
| GATGGAATAI  | GGTATTCCTT                  | CTCATTATCA     | TAAGATTTAT             | ACCATCGTAG                    | ;                   |
| TGATCTAGCT  | GGIMIICCII                  | CDTCCTTCTC     | ארב ברנבונו בר         | GCAGAAACTT                    | TTTGGGCGAG          |
| AAAACAAACG  | ATCTGATAGT                  | CCACACTCAC     | TACCTCCTAC             | TTATAAGGG                     | 1                   |
| TTATTATCCT  | ATCTGATAGI                  | CACACTOACO     | ጥር እጥጥጥ የርጥኘ           | TCCATGCAG                     | TAAGAATTTT          |
| AAACCAGCGG  | GTTCCCTAGC                  | MCMCTCATCC     | ACATCACATC             | CTGATTTGG                     | <b>A</b>            |
| ACAACTGCTG  | AAGGAGGTAG                  | TGTGACATGG     | MUNICHCHT<br>MUNICHCHT | CATGGTCAGA                    | CAAAGGATGC          |
| TGACGAAGAG  | ATGTATAAAG                  | AGTTTCAGAI     | 1 1MC1C1C11            | · CTTOOTOTO                   |                     |
| ATTAGCTAAG  | ACACAATTAG                  | GGTCATGGGA     | AIAIGACAII             | memmemeeal                    | ላ ጥጥልርልልCGTT        |
| GTTACAAGTG  | TAATATGACA                  | GATATTATGG     | CAGGTATCGG             | TCIIGIGCAA                    | A TTAGAACGTT        |
| ACCCATCTT   | GTTGAATCGT                  | CGCAGAGAAA     | TCATTGAGAA             | ATACAMIGC                     |                     |
| GGCTTTGAGG  | GGACTTCGAT                  | TAAGCCGTTG     | GTACACCTGA             | A CGGAAGAIAA                  | ACAATCGTCT          |
| * mcc* 00000  | <sup>*</sup> አመአጥሮእሮፎሮሽ     | TODACABGGC     | : TATACTTTAC           | S AACAACGAAA                  | 1                   |
| <b>ምር</b> አስርጥር <u>እ</u> ጥባ   | CAAAAAATGG                  | CTGAAGCAGG     | ; TATTGCGTG(           | ; AATGTTCAC:                  | ACAAACCATI          |
| $\mathbf{x}$ $\mathbf{c}$ $\mathbf{c}$ $\mathbf{m}$ $\mathbf{c}$ $\mathbf{m}$ $\mathbf{c}$ $\mathbf{m}$ $\mathbf{c}$ $\mathbf{c}$ | · አሮኔርርርጥልርጀ                | ι διαδητίτις   | TTTTGAAAT              | AAAGATTII                     | •                   |
| CCN NTCCCT7   | \                           | r GAAAATGAAG   | TTACACTGC              | TOTTCATAC                     | AACTIGAGIG          |
| 2 m C 2 2 C 2 M C 2   | P CCACTATCTC                | ι δυδισάδους Ι | ' TTTTAAAAA            | [ TGTTAGTAG                   | -1.                 |
| ርአመጥአርመጥል'  | r TTTGGAAGG                 | A GATATGGTGG   | S AAAGAGATA'.          | I GGIGGAAAG                   | A GACACGIIGG        |
| መጽመረመጽሞሽሽ   | P አአጥርርርርርጥርር               | : TGGAATACAG   | CTAAGTATA:             | r ATCTGAATC                   | H.                  |
| <u>አመርርአርምር</u> እ   | TGTTGGACCA                  | A AACACACCA    | A AATTGGGAAG           | TTATAATCG                     | I TGATGATIGI        |
| MULTOHOTOM  | G AAACTGAAA                 | AGTTGTTTCC     | CATTTCAAA              | G ATTCAAGAA                   | r                   |
| TCTAATGAC   | o the clother               |                |                        |                               |                     |

DNA Serotype 7

modely margues

| 8/ | 59 |  |
|----|----|--|

| ÄAAGTTTTTT | AAAAATTCGA | ATAATTTAGG | GGCAGCTCTA | ACACGAAATA | AGGCACTAAG   |
|------------|------------|------------|------------|------------|--------------|
| AAAAGCTAGA | GGTAGGTGGA | TTGCGTTCTT | GGATTCAGAT | GATTTATGGC |              |
| ACCCGAGTAA |            |            |            |            | TCATTTACT.T. |
| ATCACAATTT | TGAAAAGATT | GATGAATCTA | GTCAGTCTTT | ACGTGTCCTG |              |
| GTGTCAGGAC | CAGCAATTGT | GACTAGAAAA | ATGATGTACA | ATTACGGCTA | TCCAGGGTGT   |
| TTGACTTTCA | TGTATGATGC | AGACAAAATG | GGTTTAATTC | AGATAAAAGA |              |
| TATAAAGAAA | AATAACGATT | ATGCGATATT | ACTTCAATTG | TGTAAGAAGT | ATGACTGTTA   |
| TCTTTTAAAT | GAAAGTTTAG | CTTCGTATCG | AATTAGAAAA | AA         |              |

Fig. 6 cont.

SEQ. ID. NO. 43

modery margins

| WO-00/05378 |            | 49,        | /59 `      | $\sim U$   | PCT/NL99/00460 |
|-------------|------------|------------|------------|------------|----------------|
| AAHKHVPLME  | YNPHEAVKNN | IFGTKNVAEA | AKTAKVAKFV | MVSTDKAVNP | PNVMGATKRV     |
|             |            |            |            | KGGPVTVTDF |                |
| RMTRYFMTIP  | EASRLVIQAG | HLAKGGEIFV | LDMGEPVQIL | ELARKVILLS | GHTEEEIGIV     |
| ESGIRPGEKL  | YEELLSTEER | VSEQIHEKIF | VGRVTNKQSD | IVNSFINGLL | <b></b> ) (    |
| QKDRNELKDM  | LIEFAKQE   |            |            |            |                |

Fig. 6 cont.

CPS7E

SEQ. ID. NO. 44

modety margues

WO-00/05378
MTRVELITRE FFKKNEATSK YFQKIESRRG ELFIKFFMDK LLALILLLLL SPVIIILAIW

IKLDSKGPIF YRQERVTRYG RIFRIFKFRT MISDADKVGS LVTVGQDNRI

TKVGHIIRKY RLDEVPQLFN VLMGDMSFVG VRPEVQKYVN QYTDEMFATL LLPAGITSPA

SIAYKDEDIV LEEYCSQGYS PDEAYVQKVL PEKMKYNLEY IRNFGIISDF

Fig. 6 cont.

CPS7F

SEQ. ID. NO. 45

morlify margins

|                 |  | 1  |   |
|-----------------|--|--|---|
|                 | 51/59  |  | PCT/NL99/00460  |
| AEID EVIDTLKSGW | ITTGPKTKEL   | ERRLSVFTGT   | NKTVCLNSAT  |
| EVIV PAMTYTASCS | VITHVGATPV   | MVDIQKNSFE   | . *   |
| IIPV DLAGIPCDYD | KIYTIVENKR   | SLYVASDNKW   | QKLFGRVIIL  |
|                 |  | rameno en pot D  |   |
| TKDA LAKTOLGSWE | YDIVIPGYKC   | NMTDIMAGIG   | LVQLERYPSL  |
| GTSI KPLVHLTEDK | QSSMHLYITH   | LQGYTLEQRN   |   |
| YKPL PLLTAYKNLG | FEMKDFPNAY   | QYFENEVTLP   | LHTNLSDEDV  |
|                 |  |  |   |
|                 | AEID EVIDTLKSGW<br>EVIV PAMTYTASCS<br>IIPV DLAGIPCDYD<br>GSLA DFTSFSFHAV<br>TKDA LAKTQLGSWE<br>GTSI KPLVHLTEDK | AEID EVIDTLKSGW ITTGPKTKEL EVIV PAMTYTASCS VITHVGATPV IIPV DLAGIPCDYD KIYTIVENKR GSLA DFTSFSFHAV KNFTTAEGGS PTKDA LAKTQLGSWE YDIVIPGYKC GTSI KPLVHLTEDK QSSMHLYITH | 51/59  AEID EVIDTLKSGW ITTGPKTKEL ERRLSVFTGT EVIV PAMTYTASCS VITHVGATPV MVDIQKNSFE TIPV DLAGIPCDYD KIYTIVENKR SLYVASDNKW GSLA DFTSFSFHAV KNFTTAEGGS VTWRSHPDLD PTKDA LAKTQLGSWE YDIVIPGYKC NMTDIMAGIG GGTSI KPLVHLTEDK QSSMHLYITH LQGYTLEQRN EYKPL PLLTAYKNLG FEMKDFPNAY QYFENEVTLP |

Fig. 6 cont.

CPS7G

SEQ. ID. NO. 46

modity margins

PCT/NL99/00460

52/59

MVERDMVERD TLVSIIMPSW NTAKYISESI QSVLDQTHQN WELIIVDDCS NDETEKVVSH

FKDSRIKFFK NSNNLGAALT RNKALRKARG RWIAFLDSDD LWHPSKLEKQ

LEFMKNNGYS FTYHNFEKID ESSQSLRVLV SGPAIVTRKM MYNYGYPGCL TFMYDADKMG

LIQIKDIKKN NDYAILLQLC KKYDCYLLNE SLASYRIRK

Fig. 6 cont.

CPS7H

SEQ. ID. NO. 47

modify margans

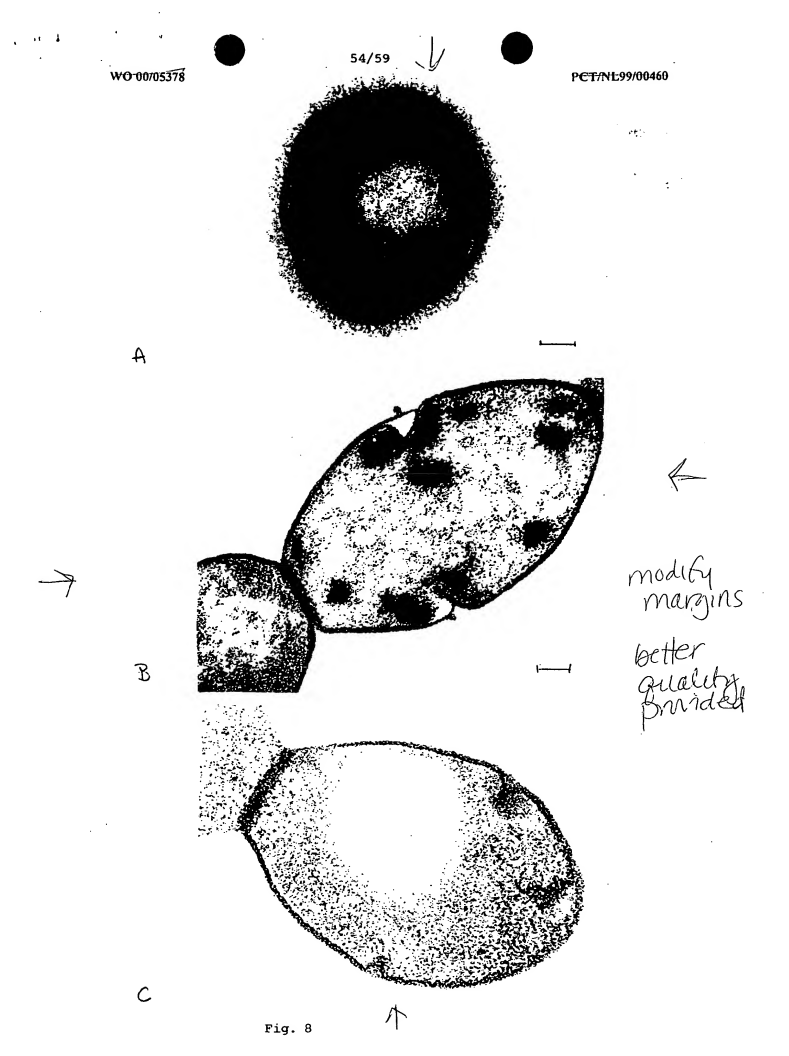
|               | -     | <br>1 1 11 1 | 11 1 1  | 1111 11 | 1 1 1 111 | EYAEQDGRIK<br>     <br>AYAKKDSRIR | 60<br>60   | Ŀ |
|---------------|-------|--------------|---------|---------|-----------|-----------------------------------|------------|---|
| $\rightarrow$ | CPCCC | <br>11111    | 1 1 1 1 | [ ]     | 1 1       | SDLSGGLLAT     NALVAVAG           | 120<br>117 |   |

Cps2J (SEQ. ID. NO. 51)

Fig. 7

Cps2K (SEQ. ID. NO. 52)

modely marzino

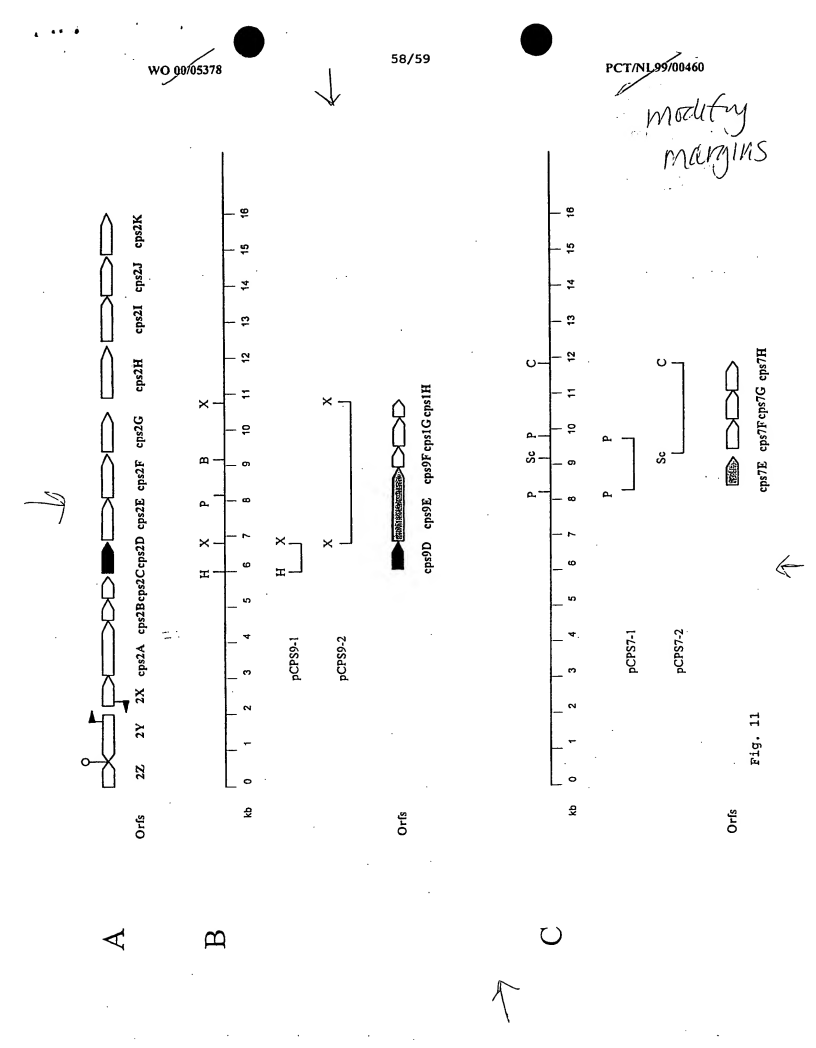


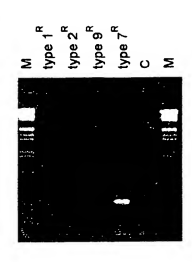
SEQ. ID. NO. 49 SEQ. ID. NO. 50 SEQ. ID. NO. 48 17084 19903 (1) 10508 AAGGGCAC (2) 16985 GGGGCAC

modify marguns

1g. 1

7





better quality copy pm/ded

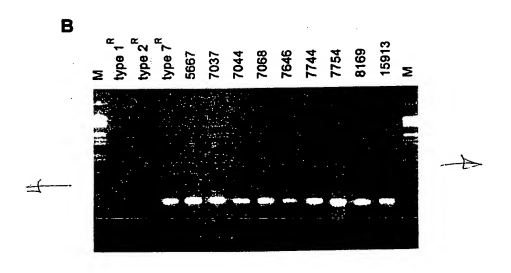


Fig. 12